

SEARCH REQUEST FORM

1.737

Requestor's
Name:

PHILLIP GAMBEL

Serial

Number:

08/487283

Date:

1/28/97

Phone:

303.3997

Art Unit:

1806

Search Topic:

Please write a detailed statement of search topic. Describe specifically as possible the subject matter to be searched. Define any terms that may have a special meaning. Give examples or relevant citations, authors, keywords, etc., if known. For sequences, please attach a copy of the sequence. You may include a copy of the broadest and/or most relevant claim(s).

SEQ / INTERFERENCE SEARCH

SEQ ID NO. 1 + 2
= =

REQUEST IN

Three

STAFF USE ONLY

Date completed:

1/28/98

Searcher:

m

Terminal time:

10

Elapsed time:

CPU time:

Total time:

10

Number of Searches:

1

Number of Databases:

5

Search Site

☐ STIC☒ CM-1☐ Pre-S

Type of Search

☐ N.A. Sequence☒ A.A. Sequence ^{x2}☐ Structure☐ Bibliographic

Vendors

☒ IG m☐ STN☐ Dialog☐ APS☐ Geninfo☐ SDC☐ DARC/Questel☐ Other


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#cross-references MUID:88209511
#accession A27689
#molecule_type mRNA
#residues 412-1676 #label WET
#cross-references GB:M18879
REFERENCE A01267
#authors Fernandez, H.N.; Hugli, T.E.
#journal J. Biol. Chem. (1978) 253:6955-6964
#title Primary structural analysis of the polypeptide portion of
human C5a anaphylatoxin. Polypeptide sequence determination
and assignment of the oligosaccharide attachment site in
C5a.
#cross-references MUID:79005687
#accession A01267
#molecule_type protein
#residues 678-751 #label FER
REFERENCE A01266
#authors Lundwall, A.B.; Wetzel, R.A.; Kristensen, T.; Whitehead,
A.S.; Woods, D.E.; Ogden, R.C.; Colten, H.R.; Tack, B.F.
#journal J. Biol. Chem. (1985) 260:2108-2112
#title Isolation and sequence analysis of a cDNA clone encoding the
fifth complement component.
#cross-references MUID:85130937
#accession A01266
#molecule_type mRNA
#residues 412-854,
'SLAISPRLECNKGTSGCHKLRPGSDSPASQVAGITGTHHQAQPT'
#label LUN
#cross-references GB:K02874
#note the carboxyl-terminal part of the sequence in this
report appears to be derived from translation of an
AUJ repeat sequence
REFERENCE S15121
#authors Bohnsack, J.F.; Mollison, K.W.; Buko, A.M.; Ashworth, J.C.;
Hill, H.R.
#journal Biochem. J. (1991) 273:635-640
#title Group B streptococci inactivate complement component C5a by
enzymic cleavage at the C-terminus.
#cross-references MUID:91144547
#contents annotation
#COMMENT Complement C5 contains two disulfide-linked chains, formed by
removal of four basic residues. C5 convertase releases C5a
anaphylatoxin from the amino end of the alpha chain, generating
C5b (beta and alpha' chains).
#COMMENT Activation of C5 initiates the spontaneous assembly of the late
complement components, C5-C9, into the membrane attack complex.
C5b has a transient binding site for C6. The C5b-C6 complex is
the foundation upon which the membrane attack complex is
assembled.
#COMMENT C5a has potent spasmogenic and chemotactic activity.
GENETICS
#gene GDB:C5
#cross-references GDB:119734
#map_position 9q33-9q33
#CLASSIFICATION #superfamily alpha-2-macroglobulin
#KEYWORDS complement alternate pathway; complement pathway; cytolysis;
glycoprotein; inflammation; membrane attack complex; plasma
FEATURE
1-18 #domain signal sequence #status predicted #label SIC\
19-673, 678-1676 #product complement C5 #status predicted #label MAT\
19-673, 752-1676 #product C5b #status predicted #label C5B\
19-673 #product complement C5 and C5b beta chain #status
predicted #label C5B8\
678-1676 #product complement C5 alpha chain #status predicted

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#label C5A\
#product C5a anaphylatoxin #status experimental #label
C5F\
#product C5b alpha' chain #status predicted #label C5BA\
678-751
752-1676
567-810, 634-669,
698-724, 699-731,
711-732, 866-1527,
1101-1159,
1375-1505,
1405-1474,
1520-1525,
1532-1606,
1553-1676,
1654-1657
741
#disulfide bonds #status predicted\
#binding site carbohydrate (Asn) (covalent) #status
experimental\
#cleavage site Arg-Leu (C5 convertase) #status
experimental\
#binding site carbohydrate (Asn) (covalent) #status
predicted
SUMMARY #length 1676 #molecular-weight 188330 #checksum 3858
Query Match 100.0%; Score 141; DB 2; Length 1676;
Best Local Similarity 100.0%; Pred. No. 3,46e-20;
Matches 21; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Db 872 vidhqtkskcvrkveqss 892
|||||
Qy 1 VIDHQTkskcvrkveqss 21
|||||
RESULT 2
ENTRY #type complete
TITLE complement C5 precursor - mouse
CONTAINS C5a anaphylatoxin; C5b
ORGANISM #formal_name Mus musculus #common_name house mouse
DATE 19-Nov-1988 #sequence_revision 15-Oct-1994 #text_change
16-Feb-1997
ACCESSIONS A35530; A27538; A40429
REFERENCE A35530
#authors Wetzel, R.A.; Fleischer, D.T.; Haviland, D.L.
#journal J. Biol. Chem. (1990) 265:2435-2440
#title Deficiency of the murine fifth complement component (C5). A
2-base pair gene deletion in a 5'-exon.
#cross-references MUID:90153853
#accession A35530
#molecule_type mRNA
#residues 1-215, 'L' #label WET
#cross-references GB:J05234
REFERENCE A27538
#authors Wetzel, R.A.; Ogata, R.T.; Tack, B.F.
#journal Biochemistry (1987) 26:737-743
#title Primary structure of the fifth component of murine
complement.
#cross-references MUID:87185363
#accession A27538
#molecule_type mRNA
#residues 'PGL', 44-1680 #label WET2
REFERENCE A40429
#authors Haviland, D.L.; Haviland, J.C.; Fleischer, D.T.; Wetzel, R.A.
#journal J. Biol. Chem. (1991) 266:11818-11825
#title Structure of the murine fifth complement component (C5) gene.
A large, highly interrupted gene with a variant donor
splice site and organizational homology with the third and

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fourth complement component genes.

#cross-references MUID:91268053
#accession A40429
#molecule_type DNA
#residues 1-15 ##label HAV
#cross-references GB:M64852
COMMENT Complement C5 contains two disulfide-linked chains, formed by removal of four basic residues. C5 convertase releases C5a anaphylatoxin from the amino end of the alpha chain, generating C5b (beta and alpha' chains).
COMMENT Activation of C5 initiates the spontaneous assembly of the late complement components, C5-C9, into the membrane attack complex. C5b has a transient binding site for C6. The C5b-C6 complex is the foundation upon which the membrane attack complex is assembled.
COMMENT C5a has potent spasmogenic and chemotactic activity.

GENETICS

#map_position 2
#introns 22/3; 86/3; 140/3; 164/3; 195/2; 223/1; 253/2; 291/3; 334/1; 372/3; 434/3; 502/3; 572/3; 622/3; 667/1; 691/1; 757/1; 787/2; 812/1; 858/3; 934/3; 955/1; 985/1; 1056/1; 1081/2; 1134/3; 1166/3; 1224/1; 1292/3; 1343/3; 1364/3; 1392/1; 1411/2; 1445/3; 1470/3; 1506/1; 1534/1; 1564/1; 1592/1; 1637/2

CLASSIFICATION

#superfamily alpha-2-macroglobulin
complement alternate pathway; complement pathway; cytolysis; glycoprotein; inflammation; membrane attack complex; plasma

FEATURE

1-18 #domain signal sequence #status predicted #label SIG
19-674, 679-1679 #product complement C5 #status predicted #label MAT
19-674, 756-1679 #product C5b #status predicted #label C5b
19-674 #product complement C5 and C5b beta chain #status predicted #label C5b
679-1679 #product complement C5 alpha chain #status predicted #label C5a
679-755 #product C5a anaphylatoxin #status predicted #label C5T
756-1679 #product C5b alpha' chain #status predicted #label C5Ba
567-814, 635-670, 702-728, 703-735, 715-736, 870-1531, 1105-1163, 1379-1509, 1409-1478, 1524-1529, 1536-1609, 1557-1679, 1657-1660 #disulfide bonds #status predicted
915, 1119, 1633 #binding_site carbohydrate (Asn) (covalent) #status predicted

SUMMARY

#length 1680 #molecular-weight 188876 #checksum 3888
Query Match 48.9%; Score 69; DB 2; Length 1680;
Best Local Similarity 47.1%; Pred. No. 9.17e-03;
Matches 8; Conservative 7; Mismatches 2; Indels 0; Gaps 0;

Db 880 hsrpvcvfrqiegcs 896

: :::::|:|:|:|:|

Qy 5 QCTKSKVKRKVEGSS 21

RESULT 3

ENTRY S76070 #type complete
TITLE hypothetical protein - Synecocystis sp. (PCC 6803)
ORGANISM #formal_name Synecocystis sp.

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#variety PCC 6803
DATE 25-Apr-1997 #sequence_revision 25-Apr-1997 #text_change 25-Apr-1997
ACCESSIONS S76070
REFERENCE S74322
#authors Kaneko, T.; Sato, S.; Kotani, H.; Tanaka, A.; Asamizu, E.; Nakamura, Y.; Miyajima, N.; Hiroseawa, M.; Sugitara, M.; Sasamoto, S.; Kimura, T.; Hosouchi, T.; Matsuno, A.; Muraki, A.; Nakazaki, N.; Nario, K.; Okumura, S.; Shimpo, S.; Takeuchi, C.; Wada, T.; Watanabe, A.; Yamada, M.; Yasuda, M.; Tabata, S.
#journal DNA Res. (1996) 3:109-136
#title Sequence analysis of the genome of the unicellular cyanobacterium Synecocystis sp. PCC6803. II. Sequence determination of the entire genome and assignment of potential protein-coding regions.

#accession

S76070 preliminary
#molecule_type DNA
#residues 1-213 ##label KAN
#cross-references EMBL:D63999
#note the nucleotide sequence was submitted to the EMBL Data Library, June 1996

SUMMARY

#length 213 #molecular-weight 22745 #checksum 2191
Query Match 44.0%; Score 62; DB 11; Length 213;
Best Local Similarity 35.0%; Pred. No. 2.43e-01;
Matches 7; Conservative 7; Mismatches 6; Indels 0; Gaps 0;

Db 92 ilayigakascikpkvss 111

:: :|:|:|:|:|

Qy 1 VIDHQGTRSKVKRKVEGSS 20

RESULT 4

ENTRY S49087 #type complete
TITLE lactoferrin binding protein - Neisseria meningitidis
ORGANISM #formal_name Neisseria meningitidis
DATE 16-Feb-1995 #sequence_revision 12-May-1995 #text_change 12-May-1995

ACCESSIONS

REFERENCE S49087
#authors Pettersson, A.M.; Klarenbeek, X.Y.Z.; van Deurzen, X.Y.Z.; Poolman, X.Y.Z.; Tomassen, X.Y.Z.

#submission submitted to the EMBL Data Library, June 1994
#description Molecular characterization of the structural gene for the lacto-ferrin receptor of the meningococcal strain H44/76.

#accession

S49087 preliminary
#status preliminary
#molecule_type DNA
#residues 1-940 ##label PET
#cross-references EMBL:X79838
SUMMARY #length 940 #molecular-weight 105347 #checksum 8194

Query Match 41.8%; Score 59; DB 10; Length 940;

Best Local Similarity 50.0%; Pred. No. 9.30e-01;

Matches 7; Conservative 5; Mismatches 2; Indels 0; Gaps 0;

Db 592 rsrkcvprkingen 605

:|:|:|:|:|

Qy 8 KSKVKRKVEGSS 21

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ENTRY          A56678      #type complete
TITLE          yemanuclein-alpha - fruit fly (Drosophila melanogaster)
ORGANISM       #formal name Drosophila melanogaster
DATE          08-Jul-1995 #sequence_revision 03-Aug-1995 #text_change
ACCESSIONS    A56678; S22146
REFERENCE     A56678
AUTHORS       Alt-Ahmed, O.; Bellon, B.; Capri, M.; Joblet, C.;
              Thomas-Delaage, M.
JOURNAL       Mech. Dev. (1992) 37:69-80
TITLE         The yemanuclein-alpha: a new Drosophila DNA binding protein
              specific for the oocyte nucleus.
ACCESSION     A56678
STATUS        #status preliminary
MOLECULE_TYPE #molecule_type DNA
RESIDUES      #residues 1-1002 #label A1E
CROSS-REFS    #cross-references GB:X63503
GENETICS
GENE          FlyBase:yem4agr
CROSS-REFS    #cross-references FlyBase:FBgn0005596
INTRONS       80/3; 154/3; 428/1 477/2; 557/2
KEYWORDS      DNA binding; oocyte
SUMMARY       #length 1002 #molecular-weight 109310 #checksum 4278

Query Match   41.1%; Score 59; DB 15; Length 1002;
Best Local Similarity 50.0%; Pred. No. 9.30e-01;
Matches 6; Conservative 5; Mismatches 1; Indels 0; Gaps 0;

Db 47 tktakirikid 58
    :::| | | | |
Qy 7 TKSSKCVQKVE 18

RESULT 6
ENTRY 149364 #type complete
TITLE protein tyrosine phosphatase - mouse
ORGANISM #formal name Mus musculus #common name house mouse
DATE 02-Jul-1996 #sequence_revision 02-Jul-1996 #text_change
ACCESSIONS 149364
REFERENCE 149364
AUTHORS Wishart, M.J.; Denu, J.M.; Williams, J.A.; Dixon, J.E.
JOURNAL J. Biol. Chem. (1995) 270:26782-26785
TITLE A single mutation converts a novel-phosphotyrosine binding
       domain into a dual-specificity phosphatase.
ACCESSION 149364
STATUS preliminary; translated from GB/EMBL/DBJ
MOLECULE_TYPE mRNA
RESIDUES 1-205 #label RES
CROSS-REFS #cross-references EMBL:U34973; NID:g1063624; CDS PID:g1063625
SUMMARY #length 205 #molecular-weight 23683 #checksum 2745

Query Match   41.1%; Score 58; DB 16; Length 205;
Best Local Similarity 25.0%; Pred. No. 1.44e+00;
Matches 5; Conservative 9; Mismatches 6; Indels 0; Gaps 0;

Db 51 ilqthgithiicirqnean 70
    :::| | | | |
Qy 1 VIDHQGTSKSCVRQKVE 20

RESULT 7
ENTRY 149365 #type complete
TITLE protein tyrosine phosphatase - mouse
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ORGANISM       #formal name Mus musculus #common name house mouse
DATE          02-Jul-1996 #sequence_revision 02-Jul-1996 #text_change
ACCESSIONS    149365
REFERENCE     149364
AUTHORS       Wishart, M.J.; Denu, J.M.; Williams, J.A.; Dixon, J.E.
JOURNAL       J. Biol. Chem. (1995) 270:26782-26785
TITLE         A single mutation converts a novel-phosphotyrosine binding
              domain into a dual-specificity phosphatase.
ACCESSION     149365
STATUS        #status preliminary; translated from GB/EMBL/DBJ
MOLECULE_TYPE #molecule_type mRNA
RESIDUES      #residues 1-223 #label RES
CROSS-REFS    #cross-references EMBL:U34973; NID:g1063624; CDS PID:g1063626
GENETICS
INTRONS       168/3
SUMMARY       #length 223 #molecular-weight 25416 #checksum 359

Query Match   41.1%; Score 58; DB 16; Length 223;
Best Local Similarity 25.0%; Pred. No. 1.44e+00;
Matches 5; Conservative 9; Mismatches 6; Indels 0; Gaps 0;

Db 51 ilqthgithiicirqnean 70
    :::| | | | |
Qy 1 VIDHQGTSKSCVRQKVE 20

RESULT 8
ENTRY 168524 #type complete
TITLE ribosomal protein L34 - human
ORGANISM #formal name Homo sapiens #common name man
DATE 24-May-1996 #sequence_revision 24-May-1996 #text_change
ACCESSIONS 168524
REFERENCE 154209
AUTHORS Rommens, J.M.; Durocher, F.; McArthur, J.; Tonin, P.;
        Leblanc, J.
JOURNAL Genomics (1995) 28:530-542
TITLE Generation of a transcription map at the HSD17B locus
        centromeric to BRCA1 at 17q21.
ACCESSION 168524
STATUS preliminary; translated from GB/EMBL/DBJ
MOLECULE_TYPE mRNA
RESIDUES 1-117 #label RES
CROSS-REFS #cross-references GB:L38941; NID:g1008855; CDS PID:g1008856
CLASSIFICATION #superfamily rat ribosomal protein L34
SUMMARY #length 117 #molecular-weight 13305 #checksum 4392

Query Match   40.4%; Score 57; DB 8; Length 117;
Best Local Similarity 41.7%; Pred. No. 2.22e+00;
Matches 5; Conservative 6; Mismatches 1; Indels 0; Gaps 0;

Db 80 gmcakcvdrri 91
    :::| | | | |
Qy 6 GTKSSKCVQKRV 17

RESULT 9
ENTRY S28969 #type complete
TITLE N-carbamoylsarcosine amidohydrolase (EC 3.5.1.59) -
        Arthrobacter sp.
ORGANISM #formal name Arthrobacter sp.
DATE 19-Mar-1997 #sequence_revision 19-Mar-1997 #text_change
```

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ACCESSIONS 528969
REFERENCE
#authors Romao, M.J.; Turk, D.; Comis-Rueth, F.X.; Huber, R.;
Schumacher, G.; Moellerling, H.; Ruessmann, L.
#journal J. Mol. Biol. (1992) 226:1111-1130
#title Crystal structure analysis, refinement and enzymatic reaction
mechanism of N-carbamoylserine amidohydrolase from
Arthrobacter sp. at 2.0 A resolution.
#accession 528969
#status preliminary
#residues 1-264 #label ROM
SUMMARY #length 264 #molecular-weight 29057 #checksum 6729

Query Match 40.4%; Score 57; DB 18; Length 264;
Best Local Similarity 40.0%; Pred. No. 2.22e+00;
Matches 6; Conservative 5; Mismatches 4; Indels 0; Gaps 0;

Db 171 gataagcvtvhtveda 185
| : : ||| : || :
Qy 6 GTKSKCVRQKVEGS 20

RESULT 10
ENTRY B33485 #type complete
TITLE spore coat protein SP70 - slime mold (Dictyostelium
discoideum)
ORGANISM #formal name Dictyostelium discoideum
DATE 09-Mar-1990 #sequence_revision 11-Sep-1992 #text_change
30-Sep-1993
ACCESSIONS B33485
REFERENCE A33485
#authors Foenbaugh, K.L.; Loomis, W.F.
#journal Mol. Cell. Biol. (1989) 9:5215-5218
#title Spore coat genes SP60 and SP70 of Dictyostelium discoideum.
#cross-references MUID:90097939
#accession B33485
#status preliminary
#molecule_type DNA; mRNA
#residues 1-537 #label FOS
#cross-references GB:M26238
#note the authors translated the codon AAT for residue 281 as
Asp
CLASSIFICATION #superfamily LDL receptor ligand-binding repeat homology
SUMMARY #length 537 #molecular-weight 56650 #checksum 2250

Query Match 39.7%; Score 56; DB 13; Length 537;
Best Local Similarity 54.5%; Pred. No. 3.42e+00;
Matches 6; Conservative 4; Mismatches 1; Indels 0; Gaps 0;

Db 291 kncgirdkve 301
| : : ||| : |||
Qy 8 KSSKCVQRKVE 18

RESULT 11
ENTRY S21825 #type complete
TITLE vicilin-like storage protein G1b1-S, embryo - maize
ORGANISM #formal name Zea mays #common_name maize
DATE 20-Feb-1995 #sequence_revision 20-Feb-1995 #text_change
20-Feb-1995
ACCESSIONS S21825
REFERENCE S21823
#authors Kriz, A.L.
#submission submitted to the EMBL Data Library, April 1991
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```
#accession S21825
#status preliminary
#molecule_type DNA
#residues 1-540 #label KRI
#cross-references EMBL:X59084
GENETICS
#gene G1b1-S
#introns 170/1; 195/2; 222/2; 319/2
SUMMARY #length 540 #molecular-weight 60239 #checksum 1419

Query Match 39.7%; Score 56; DB 13; Length 540;
Best Local Similarity 58.3%; Pred. No. 3.42e+00;
Matches 7; Conservative 3; Mismatches 2; Indels 0; Gaps 0;

Db 32 hqghkagrcvrr 43
| | | : ||| :
Qy 4 HQGTSKSCVRQ 15

RESULT 12
ENTRY A53234 #type complete
TITLE globulin-1S, G1B1S - maize
ORGANISM #formal name Zea mays #common_name maize
DATE 02-May-1994 #sequence_revision 18-Nov-1994 #text_change
05-Apr-1995
ACCESSIONS A53234
REFERENCE A53234
#authors Belanger, F.C.; Kriz, A.L.
#journal Genetics (1991) 129:863-872
#title Molecular basis for allelic polymorphism of the maize
Globulin-1 gene.
#cross-references MUID:92090707
#accession A53234
#status preliminary
#molecule_type DNA
#residues 1-573 #label BEL
#cross-references NCBI:71280; NCBI:P:71284
#experimental_source inbred line Va 26
#note sequence extracted from NCBI backbone
SUMMARY #length 573 #molecular-weight 65075 #checksum 3569

Query Match 39.7%; Score 56; DB 13; Length 573;
Best Local Similarity 58.3%; Pred. No. 3.42e+00;
Matches 7; Conservative 3; Mismatches 2; Indels 0; Gaps 0;

Db 32 hqghkagrcvrr 43
| | | : ||| :
Qy 4 HQGTSKSCVRQ 15

RESULT 13
ENTRY A32494 #type complete
TITLE transposable element Txc protein 1 - African clawed frog
ORGANISM #formal name Xenopus laevis #common_name African clawed frog
DATE 12-Oct-1989 #sequence_revision 31-Dec-1993 #text_change
31-Dec-1993
ACCESSIONS A32494
REFERENCE A32494
#authors Garrett, J.E.; Knutzon, D.S.; Carroll, D.
#journal Mol. Cell. Biol. (1989) 9:3018-3027
#title Composite transposable elements in the Xenopus laevis genome.
#cross-references MUID:89384562
#accession A32494
#status preliminary
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```
#molecule type DNA
##residues 1-775 ##label GAR
##cross-references GB:M26915
##note the authors translated the codon ATT for residue as Gln,
and AAG for residue 288 as Leu
SUMMARY #length 775 #molecular-weight 82355 #checksum 6734

Query Match 39.0%; Score 55; DB 15; Length 775;
Best Local Similarity 46.7%; Pred. No. 5.22e+00;
Matches 7; Conservative 5; Mismatches 3; Indels 0; Gaps 0;

Db 617 ntsskcvsevgtp 631
Qy 7 TKSSKCVQRKVEGSS 21
:::||||| |||::
```

```
RESULT 14
ENTRY EA7119 #type complete
TITLE spore coat peptide CotZ - Bacillus subtilis
ORGANISM #formal_name Bacillus subtilis
DATE 21-Sep-1993 #sequence_revision 18-Nov-1994 #text_change
18-Nov-1994
ACCESSIONS EA7119
REFERENCE A47119
authors Zhang, J.; Fitz-James, P.C.; Aronson, A.I.
#journal J. Bacteriol. (1993) 175:3757-3766
#title Cloning and characterization of a cluster of genes encoding
polypeptides present in the insoluble fraction of the spore
coat of Bacillus subtilis.
#cross-references MUID:93285989
#accession EA7119
##status preliminary
##molecule type nucleic acid
##residues 1-148 ##label ZHA
##cross-references NCBIN:133538; NCIP:133548
##note sequence extracted from NCBI backbone
SUMMARY #length 148 #molecular-weight 16534 #checksum 4681
```

```
Query Match 38.3%; Score 54; DB 12; Length 148;
Best Local Similarity 63.6%; Pred. No. 7.93e+00;
Matches 7; Conservative 2; Mismatches 2; Indels 0; Gaps 0;

Db 4 ktsscvreave 14
I:I |||: ||
Qy 8 KSSKCVQRKVE 18
```

```
RESULT 15
ENTRY G64383 #type complete
TITLE riboflavin-specific deaminase (EC 3.5.4.-) - Methanococcus
jannaschii
ORGANISM #formal_name Methanococcus jannaschii
DATE 13-Sep-1996 #sequence_revision 13-Sep-1996 #text_change
13-Sep-1996
ACCESSIONS G64383
REFERENCE A64300
authors Bult, C.J.; White, O.; Olsen, G.J.; Zhou, L.; Fleischmann,
R.D.; Sutton, G.G.; Blake, J.A.; FitzGerald, L.M.; Clayton,
R.A.; Gocayne, J.D.; Kerlavage, A.R.; Dougherty, B.A.;
Tomb, J.F.; Adams, M.D.; Reich, C.I.; Overbeek, R.;
Kirkness, E.F.; Weinstock, K.G.; Merrick, J.M.; Glodek, A.;
Scott, J.L.; Geoghagan, N.S.M.; Weidman, J.F.; Fuhrmann,
J.L.; Nguyen, D.; Utterback, T.R.; Kelley, J.M.; Peterson,
J.D.; Sadow, P.W.; Hanna, M.C.; Cotton, M.D.; Roberts,
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K.M.; Hurst, M.A.; Kaine, B.P.; Borodovsky, M.; Klenk,
H.P.; Fraser, C.M.; Smith, H.O.; Woese, C.R.; Venter, J.C.
#journal Science (1996) 273:1058-1073
#title Complete genome sequence of the methanogenic archaeon,
Methanococcus jannaschii.
#accession G64383
##status preliminary; nucleic acid sequence not shown;
translation not shown
##molecule type DNA
##residues 1-224 ##label BUL
##cross-references GB:L7117; TIGR:MJ0671; CDS_PID:g1510756
GENETICS
#map_position REV597638-596964
#start_codon TTC
KEYWORDS hydrolase
SUMMARY #length 224 #molecular-weight 25037 #checksum 2215
```

```
Query Match 38.3%; Score 54; DB 12; Length 224;
Best Local Similarity 33.3%; Pred. No. 7.93e+00;
Matches 6; Conservative 4; Mismatches 8; Indels 0; Gaps 0;

Db 118 iledmgvervkcgrgkvd 135
::: | |||::
Qy 1 VIDHQTGKSSKCVQRKVE 18
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Search completed: Wed Jan 28 12:10:33 1998
Job time : 16 secs.

[4] RP SEQUENCE OF 678-751.
 RX MEDLINE: 79005687.
 RA FERNANDEZ H.N., HUGLI T.E.;
 RL J. BIOL. CHEM. 253:6955-6964 (1978).
 [5] RP SEQUENCE OF 678-751 FROM N.A.
 RX MEDLINE: 91144547.
 RA BOHSACK J.F., MOLLISON K.W., BUKO A.M., ASHWORTH J.C., HILL H.R.;
 RL BIOCHEM. J. 273:635-640 (1991).
 [6] RP STRUCTURE BY NMR OF C5A.
 RX MEDLINE: 88309754.
 RA ZUIDERWEG E.R., MOLLISON K.W., HENKIN J., CARTER G.W.;
 RL BIOCHEMISTRY 27:3568-3580 (1988).
 [7] RP STRUCTURE BY NMR OF C5A.
 RX MEDLINE: 89207527.
 RA ZUIDERWEG E.R., NETTESHEIM D.G., MOLLISON K.W., CARTER G.W.;
 RL BIOCHEMISTRY 28:172-185 (1989).
 [8] RP STRUCTURE BY NMR OF C5A.
 RX MEDLINE: 89274164.
 RA ZUIDERWEG E.R., FESIK S.W.;
 RL BIOCHEMISTRY 28:2387-2391 (1989).
 CC -!- FUNCTION: ACTIVATION OF C5 BY A C5 CONVERTASE INITIATES THE
 CC SPONTANEOUS ASSEMBLY OF THE LATE COMPLEMENT COMPONENTS, C5-C9,
 CC INTO THE MEMBRANE ATTACK COMPLEX. C5B HAS A TRANSIENT BINDING SITE
 CC FOR C6. THE C5B-C6 COMPLEX IS THE FOUNDATION UPON WHICH THE LYTC
 CC COMPLEX IS ASSEMBLED.
 CC -!- SUBUNIT: C5 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 BASIC
 CC RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY A DISULFIDE
 CC BOND. C5 CONVERTASE ACTIVATES C5 BY CLEAVING THE ALPHA CHAIN,
 CC RELEASING C5A ANAPHYLATOXIN & GENERATING C5B (BETA CHAIN + ALPHA
 CC CHAIN).
 CC -!- SIMILARITY: TO C3, C4 AND ALPHA-2-MACROGLOBULIN.
 CC -!- FUNCTION: DERIVED FROM PROTEOLYTIC DEGRADATION OF COMPLEMENT C5,
 CC C5 ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS. IT
 CC INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR
 CC PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND
 CC BASOPHILIC LEUKOCYTES. C5A ALSO STIMULATES THE LOCOMOTION OF
 CC POLYMORPHONUCLEAR LEUKOCYTES (CHEMOKINESIS) AND DIRECT THEIR
 CC MIGRATION TOWARD SITES OF INFLAMMATION (CHEMOTAXIS).
 CC -!- CAUTION: REF. 3 SEQUENCE DIFFERS FROM THAT SHOWN FROM POSITION 855
 CC ONWARD DUE TO THE PRESENCE OF AN ALU REPEAT.
 CC -!- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN.
 DR EMBL: M57729; G179983; --
 DR EMBL: M65134; G179692; --
 DR PIR: A40075; CSHU.
 DR PIR: S15121; S15121.
 DR HSP: P01032; IC5A.
 DR MIM: 120900; --
 DR PROSITE: PS00477; ALPHA 2 MACROGLOBULIN.
 KW COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; GLYCOPROTEIN;
 KW PLASMA; MEMBRANE ATTACK COMPLEX; CYTOLYSIS; INFLAMMATORY RESPONSE;
 KW SIGNAL; POLYMORPHISM.
 FT SIGNAL 1 18 POTENTIAL.
 FT CHAIN 19 673 COMPLEMENT C5 BETA CHAIN.
 FT PROPEP 674 677
 FT CHAIN 678 1676 COMPLEMENT C5 ALPHA CHAIN.
 FT PEPTIDE 678 751 C5A ANAPHYLATOXIN.
 FT CHAIN 752 1676 C5B (ALPHA').
 FT DOMAIN 698 732 ANAPHYLATOXIN-LIKE.
 FT DISULFID 698 724

FT DISULFID 699 731
 FT DISULFID 711 732
 FT CARBOHYD 741 741
 FT CARBOHYD 911 911 POTENTIAL.
 FT CARBOHYD 1115 1115 POTENTIAL.
 FT CARBOHYD 1630 1630 POTENTIAL.
 FT VARIANT 518 518 F -> S.
 SQ SEQUENCE 1676 AA; 188331 MW; 9D5C6E59 CRC32;
 Query Match 100.0%; Score 141; DB 2; Length 1676;
 Best Local Similarity 100.0%; Pred. No. 1.16e-24;
 Matches 21; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 Db 872 vidhgtkaskcvrkvqvegs 892
 |||||
 Qy 1 VIDHGTKSKCKVQKVEGS 21
 RESULT 2
 ID C05 MOUSE STANDARD; PRT; 1680 AA.
 AC P06684;
 DT 01-JAN-1988 (REL. 06, CREATED)
 DT 01-NOV-1990 (REL. 16, LAST SEQUENCE UPDATE)
 DT 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE)
 DE COMPLEMENT C5 PRECURSOR (CONTAINS: C5A ANAPHYLATOXIN).
 GN C5.
 OS MUS MUSCULUS (MOUSE).
 OC EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA;
 OC EUTHERIA; RODENTIA.
 RN [1]
 RP SEQUENCE FROM N.A.
 RX MEDLINE: 90153853.
 RA WETSEL R.A., FLEISCHER D.T., HAVILAND D.L.;
 RL J. BIOL. CHEM. 265:2435-2440 (1990).
 RN [2]
 RP SEQUENCE OF 41-1680 FROM N.A.
 RX MEDLINE: 87185363.
 RA WETSEL R.A., OGATA R.T., TACK B.F.;
 RL BIOCHEMISTRY 26:737-743 (1987).
 CC -!- FUNCTION: ACTIVATION OF C5 BY A C5 CONVERTASE INITIATES THE
 CC SPONTANEOUS ASSEMBLY OF THE LATE COMPLEMENT COMPONENTS, C5-C9,
 CC INTO THE MEMBRANE ATTACK COMPLEX. C5B HAS A TRANSIENT BINDING SITE
 CC FOR C6. THE C5B-C6 COMPLEX IS THE FOUNDATION UPON WHICH THE LYTC
 CC COMPLEX IS ASSEMBLED.
 CC -!- SUBUNIT: C5 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 BASIC
 CC RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY A DISULFIDE
 CC BOND. C5 CONVERTASE ACTIVATES C5 BY CLEAVING THE ALPHA CHAIN,
 CC RELEASING C5A ANAPHYLATOXIN & GENERATING C5B (BETA CHAIN + ALPHA
 CC CHAIN).
 CC -!- SIMILARITY: TO C3, C4 AND ALPHA-2-MACROGLOBULIN.
 CC -!- FUNCTION: DERIVED FROM PROTEOLYTIC DEGRADATION OF COMPLEMENT C5,
 CC C5 ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS. IT
 CC INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR
 CC PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND
 CC BASOPHILIC LEUKOCYTES. C5A ALSO STIMULATES THE LOCOMOTION OF
 CC POLYMORPHONUCLEAR LEUKOCYTES (CHEMOKINESIS) AND DIRECT THEIR
 CC MIGRATION TOWARD SITES OF INFLAMMATION (CHEMOTAXIS).
 CC -!- CAUTION: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN.
 DR EMBL: M35525; G309124; --
 DR EMBL: M35526; G309123; --
 DR PIR: A27538; A27538.
 DR PIR: A35530; A35530.
 DR HSP: P01032; IC5A.
 DR PROSITE: PS00477; ALPHA_2_MACROGLOBULIN.

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KW COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; GLYCOPROTEIN;
KW PLASMA; MEMBRANE ATTACK COMPLEX; CYTOLYSIS; INFLAMMATORY RESPONSE;
KW SIGNAL.
FT SIGNAL 1 18
FT CHAIN 19 1680
FT CHAIN 19 674
FT PROPEP 675 678
FT CHAIN 679 1680
FT PEPTIDE 679 755
FT CHAIN 756 1680
FT DOMAIN 702 736
FT DISULFID 702 728
FT DISULFID 703 735
FT DISULFID 715 736
FT CARBOHYD 427 427
FT CARBOHYD 915 915
FT CARBOHYD 1119 1119
FT CARBOHYD 1633 1633
FT VARIANT 216 216
FT VARIANT 217 1680
SQ SEQUENCE 1680 AA; 188877 MW; AA17044B CRC32;

Query Match 48.9%; Score 69; DB 2; Length 1680;
Best Local Similarity 47.1%; Pred. No. 7.22e-04;
Matches 8; Conservative 7; Mismatches 2; Indels 0; Gaps 0;

Db 880 hterparcvfgriegas 896
: ::::|::|::|::|::|
Qy 5 QGTSSKCVRKVEGSS 21

RESULT 3
ID IROA NEIME STANDARD; PRT; 943 AA.
AC Q06379;
DT 01-NOV-1995 (REL. 32, CREATED)
DT 01-NOV-1995 (REL. 32, LAST SEQUENCE UPDATE)
DT 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE)
DE IRON-REGULATED OUTER MEMBRANE PROTEIN A PRECURSOR.
GN IROA.
OS NEISSERIA MENINGITIDIS.
OC PROKARYOTA; GRACILICUTES; SCOTOBACTERIA; AEROBIC RODS AND COCCI;
OC NEISSERIACEAE.
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=BNVC;
RX MEDLINE; 94011384.
RA PETERSSON A., VAN DER LEY P., POOLMAN J.T., TOMMASSEN J.;
RL INFECT. IMMUN. 61:4724-4733(1993).
CC -!- FUNCTION: UNKNOWN. MAY BE AN IRON-SIDEROPHORE RECEPTOR.
CC -!- SUBCELLULAR LOCATION: OUTER MEMBRANE.
CC -!- INDUCTION: BY IRON-STARVATION CONDITIONS.
CC -!- SIMILARITY: LOCAL TO OTHER TONB-DEPENDENT RECEPTOR PROTEINS.
DR EMBL; X69214; G45064; -.
DR PROSITE; PS00430; TONB DEPENDENT REC 1.
DR PROSITE; PS01156; TONB DEPENDENT REC 2.
KW OUTER MEMBRANE; IRON TRANSPORT; TONB BOX; SIGNAL; RECEPTOR.
FT SIGNAL 1 27
FT CHAIN 28 943
FT CHAIN 826 943
FT SIMILAR TONB C-TERMINAL BOX.
SQ SEQUENCE 943 AA; 105424 MW; 16644948 CRC32;

Query Match 41.8%; Score 59; DB 5; Length 943;
Best Local Similarity 50.0%; Pred. No. 1.65e-01;
Matches 7; Conservative 5; Mismatches 2; Indels 0; Gaps 0;

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Db 595 tsrkcvprkingns 608
:|::|::|::|::|::|::|
Qy 8 KSSKCVRKVEGSS 21

RESULT 4
ID YEMA DROME STANDARD; PRT; 1002 AA.
AC P25992;
DT 01-MAY-1992 (REL. 22, CREATED)
DT 01-MAY-1992 (REL. 22, LAST SEQUENCE UPDATE)
DT 01-FEB-1995 (REL. 31, LAST ANNOTATION UPDATE)
DE YEMANUCLEIN-ALPHA.
GN YEMA OR YG4.5.
OS DROSOPHILA MELANOGASTER (FRUIT FLY).
OC EUKARYOTA; METAZOA; ARTHROPODA; INSECTA; DIPTERA.
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=CANTON-S;
RX MEDLINE; 92297435.
RA AIT-AHMED O., BELLON B., CAPRI M., JOBLET C., THOMAS-DELAAGE M.;
RL MECH. DEV. 37:69-80(1992).
CC -!- FUNCTION: MAY PLAY A KEY ROLE IN EGG ORGANIZATION. IT MAY BE A TRANSCRIPTIONAL REGULATOR.
CC -!- PTM: THE N-TERMINAL IS BLOCKED.
CC -!- TISSUE SPECIFICITY: OOCYTE-SPECIFIC.
CC -!- DEVELOPMENTAL STAGE: EXPRESSED AT ALL OOGENIC STAGES.
CC -!- SUBCELLULAR LOCATION: NUCLEAR.
DR EMBL; X63503; G8838; -.
DR PIR; S22146; S22146.
DR FLYBASE; FBGN005596; YEM-ALPHA.
KW NUCLEAR PROTEIN; DNA-BINDING; REPEAT.
FT DOMAIN 80 85
FT DOMAIN 207 217
FT DOMAIN 219 261
FT DOMAIN 230 253
FT REPEAT 230 241
FT REPEAT 242 253
FT VARIANT 698 698
SQ SEQUENCE 1002 AA; 109310 MW; 955FD2C1 CRC32;

Query Match 41.8%; Score 59; DB 11; Length 1002;
Best Local Similarity 50.0%; Pred. No. 1.65e-01;
Matches 6; Conservative 5; Mismatches 1; Indels 0; Gaps 0;

Db 47 tktakcirikid 58
|::|::|::|::|::|
Qy 7 TKSSKCVRKVE 18

RESULT 5
ID RL34 HUMAN STANDARD; PRT; 116 AA.
AC P49207;
DT 01-FEB-1996 (REL. 33, CREATED)
DT 01-FEB-1996 (REL. 33, LAST SEQUENCE UPDATE)
DT 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE)
DE 60S RIBOSOMAL PROTEIN L34.
GN RPL34.
OS HOMO SAPIENS (HUMAN).
OC EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA;
OC EUTHERIA; PRIMATES.
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE-OVARY;

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RX MEDLINE; 96039267.
RA ROMMENS J.M., DUROCHER F., MCARTHUR J., TONIN P., LEBLANC J.F.,
RA ALLEN T., SAMSON C., FERRI L., NAROD S., MORGAN K., SIMARD J.;
RL GENOMICS 28:530-542(1995).
CC -!- SIMILARITY: BELONGS TO THE L34E FAMILY OF RIBOSOMAL PROTEINS.
DR EMBL; L38941; G1008856; -.
KW RIBOSOMAL PROTEIN.
FT INIT MET 0 0 BY SIMILARITY.
SQ SEQUENCE 116 AA; 13174 MW; 490F4AF1 CRC32;

Query Match 40.4%; Score 57; DB 8; Length 116;
Best Local Similarity 41.7%; Pred. No. 4.57e-01;
Matches 5; Conservative 6; Mismatches 1; Indels 0; Gaps 0;

Db 79 gmcakcvrdri 90
I:::|||||::: 17
Qy 6 GTKSSKCVQRKV 17

RESULT 6
ID GSH ARTSP STANDARD; PRT; 264 AA.
AC P32400;
DT 01-OCT-1993 (REL. 27, CREATED)
DT 01-OCT-1993 (REL. 27, LAST SEQUENCE UPDATE)
DT 01-FEB-1995 (REL. 31, LAST ANNOTATION UPDATE)
DE N-CARBAMOYL-SARCOSE AMIDASE (EC 3.5.1.59) (N-CARBAMOYL-SARCOSE
DE AMIDOHYDROLASE) (CSHASE).
OS ARTHROBACTER SP.
OC PROKARYOTA; FIRMICUTES; IRREGULAR ASPOGENOUS RODS; CORYNEFORM GROUP.
RN [1]
RP X-RAY CRYSTALLOGRAPHY (2.0 ANGSTROMS), AND REVISIONS TO 184 AND 232.
RX MEDLINE; 92389321.
RA ROMAO M.J., TURK D., GOMIS-RUETH F.-X., HUBER R.;
RL J. MOL. BIOL. 226:1111-1130(1992).
CC -!- CATALYTIC ACTIVITY: N-CARBAMOYL-SARCOSE + H(2)O = SARCOSE +
CC CO(2) + NH(3).
CC -!- SUBUNIT: HOMOTETRAMER.
CC -!- COFACTOR: ONE SULFATE ION PER SUBUNIT.
CC -!- PATHWAY: DEGRADATION OF CREATININE TO GLYCINE.
DR PIR; S28969; S28969.
DR PDB; 1NBA; 22-JUN-94.
KW HYDROLASE; 3D-STRUCTURE.
FT ACT SITE 177 177 INVOLVED IN HYDROLYSIS OF THE SUBSTRATE.
SQ SEQUENCE 264 AA; 29057 MW; 81A56865 CRC32;

Query Match 40.4%; Score 57; DB 2; Length 264;
Best Local Similarity 40.0%; Pred. No. 4.57e-01;
Matches 6; Conservative 5; Mismatches 4; Indels 0; Gaps 0;

Db 171 gataagcvrhtveda 185
I:::||||: 20
Qy 6 GTKSSKCVQRKVEGS 20

RESULT 7
ID SP70 DICDI STANDARD; PRT; 537 AA.
AC P15269; P08126;
DT 01-AUG-1988 (REL. 08, CREATED)
DT 01-APR-1990 (REL. 14, LAST SEQUENCE UPDATE)
DT 01-FEB-1994 (REL. 28, LAST ANNOTATION UPDATE)
DE SPORE COAT PROTEIN SP70 PRECURSOR (BEEJIN PROTEIN).
GN COYB.
OS DICTYOSTELIUM DISCOIDEUM (SLIME MOLD).
CC EUKARYOTA; PROTOZOA; SARCOMASTIGOPHORA; SARCODINA; RHIZOPODA;

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OC EUMYCETOZOA; DICTYOSTELIA.
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE; 90097939.
RA FOSNAUGH K.L., LOOMIS W.F.;
RL MOL. CELL. BIOL. 9:5215-5218(1989).
RN [2]
RP PRELIMINARY SEQUENCE OF 72-170 FROM N.A.
RX MEDLINE; 87057653.
RA GOMER R.H., DATTA S., FIRTEL R.A.;
RL J. CELL BIOL. 103:1999-2015(1986).
DR EMBL; M26238; G167889; -.
DR PIR; B33485; B33485.
DR PIR; B25439; B25439.
DR DICTYDB; DD03009; COYB.
KW GLYCOPROTEIN; PHOSPHORYLATION; REPEAT; SPOULATION; SIGNAL.
FT SIGNAL 1 20
FT CHAIN 21 537 SPORE COAT PROTEIN SP70.
FT DOMAIN 182 250 SER/THR-RICH.
FT DOMAIN 190 248 5.5 X 11 AA TANDEM REPEATS.
FT REPEAT 190 200 1.
FT REPEAT 191 211 2.
FT REPEAT 212 222 3.
FT REPEAT 223 233 4.
FT REPEAT 234 244 5.
FT REPEAT 245 248 6 (INCOMPLETE).
FT REPEAT 256 263 PRESPORE MOTIF.
FT REPEAT 284 291 PRESPORE MOTIF.
FT REPEAT 364 371 PRESPORE MOTIF.
FT CARBOHYD 97
SQ SEQUENCE 537 AA; 56650 MW; 5D59CBAC CRC32;

Query Match 39.7%; Score 56; DB 9; Length 537;
Best Local Similarity 54.5%; Pred. No. 7.54e-01;
Matches 6; Conservative 4; Mismatches 1; Indels 0; Gaps 0;

Db 291 kngecirldkve 301
I:::|:|:| 18
Qy 8 KSSKCVQRKVE 18

RESULT 8
ID GLB1 MAIZE STANDARD; PRT; 573 AA.
AC P15590;
DT 01-APR-1990 (REL. 14, CREATED)
DT 01-AUG-1990 (REL. 15, LAST SEQUENCE UPDATE)
DT 01-OCT-1994 (REL. 30, LAST ANNOTATION UPDATE)
DE GLOBULIN-1 S ALLELE PRECURSOR (GLB1-S) (7S-LIKE).
GN GLB1.
OS ZEA MAYS (MAIZE).
OC EUKARYOTA; PLANTA; EMBRYOPHYTA; ANGIOSPERMAE; MONOCOTYLEDONEAE;
OC CYPERALES; GRAMINEAE.
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN-CV. INBRED LINE VA26;
RA BELANGER F.C., KRIZ A.L.;
RL PLANT PHYSIOL. 91:636-643(1989).
RN [2]
RP SEQUENCE OF 87-100.
RX MEDLINE; 89374022.
RA KRIZ A.L.;
RL BIOCHEM. GENET. 27:239-251(1989).
CC -!- SIMILARITY: TO OTHER 7S SEED STORAGE PROTEINS (PHASEOLIN, VICILIN,
CC CONVICILIN, CONGLYCININ, ETC.).

CC -!- POLYMORPHISM: THE THREE MOST COMMONLY OCCURRING GIBI ALLELES HAVE
CC THE DESIGNATION L, I, AND S FOR LARGE, INTERMEDIATE, AND SMALL
CC PROTEINS, RESPECTIVELY.
CC -!- PTM: THREE PROTEIN-PROCESSING STEPS OCCUR IN THE FORMATION OF THE
CC MATURE PROTEIN FROM THE PRIMARY TRANSLATION PRODUCT.

DR EMBL; M24845; G168481; -.

DR HSP; P02853; 1CAU.

DR MAIZEDB; 30181; -.

KW SEED STORAGE PROTEIN; SIGNAL.

FT SIGNAL 1 18 OR 21 (POTENTIAL).

FT PROPEP 19 86

FT CHAIN 87 573 GLOBULIN-1 S.

FT CARBOHYD 349 349 POTENTIAL.

SQ SEQUENCE 573 AA; 65029 MW; 7E755E20 CRC32;

Query Match 39.7%; Score 56; DB 4; Length 573;

Best Local Similarity 58.3%; Pred. No. 7.54e-01;

Matches 7; Conservative 3; Mismatches 2; Indels 0; Gaps 0;

Db 32 hghkgcgcrr 43

|||||:

Qy 4 HQTGKSKVRQ 15

RESULT 9

ID RPOB PSEPU STANDARD; PRT; 1357 AA.

AC P19175;

DT 01-NOV-1990 (REL. 16, CREATED)

DT 01-MAR-1992 (REL. 21, LAST SEQUENCE UPDATE)

DT 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE)

DE DNA-DIRECTED RNA POLYMERASE BETA CHAIN (EC 2.7.7.6) (TRANSCRIPTASE

DE BETA CHAIN) (RNA POLYMERASE BETA SUBUNIT).

GN RPOB.

OS PSEUDOMONAS PUTIDA.

OC PROKARYOTA; GRACILICUTES; SCOTOBACTERIA; AEROBIC RODS AND COCCI;

OC PSEUDOMONADACEAE.

RN [1]

RP SEQUENCE FROM N.A.

RA BORODIN A.M., DANILKOVICH A.V., ALLIKMETS R.L., ROSTAPSHOV V.M.,

RA CHERNOV I.P., AZHIKINA T.L., MONASTYRSKAYA S., SVERDLOV D.;

RL DOKL. BIOCHEM. 302:1261-1265(1988).

RN [2]

RP SEQUENCE OF 1036-1357 FROM N.A.

RX MEDLINE; 89117617.

RA BORODIN A.M., DANILKOVICH A.V., CHERNOV I.I., AZHIKINA T.L.,

RA ROSTAPSHOV V.M., MONASTYRSKAYA G.S.;

RL BIOORG. KHIM. 14:1179-1182(1988).

CC -!- FUNCTION: DNA-DEPENDENT RNA POLYMERASE CATALYZES THE TRANSCRIPTION

CC OF DNA INTO RNA USING THE FOUR RIBONUCLEOSIDE TRIPHOSPHATES AS

CC SUBSTRATES.

CC -!- CATALYTIC ACTIVITY: N NUCLEOSIDE TRIPHOSPHATE = N PYROPHOSPHATE +

CC RNA (N).

CC -!- SUBUNIT: THE ENZYME CONSISTS OF THE SIGMA CHAIN AND THE CORE

CC ENZYME WHICH IS COMPOSED OF 2 ALPHA CHAINS, 1 BETA CHAIN, AND 1

CC BETA' CHAIN.

CC -!- SIMILARITY: BELONGS TO THE RNA POLYMERASE BETA CHAIN FAMILY.

DR EMBL; X15849; G45729; -.

DR EMBL; M38319; G151547; -.

KW TRANSCRIPTION; DNA-DIRECTED RNA POLYMERASE.

FT CONFLICT 1180 1180 T -> N (IN REF. 2).

FT CONFLICT 1184 1184 I -> V (IN REF. 2).

FT CONFLICT 1236 1236 F -> S (IN REF. 2).

SQ SEQUENCE 1357 AA; 151305 MW; BBF88A37 CRC32;

Query Match 39.7%; Score 56; DB 8; Length 1357;
Best Local Similarity 41.2%; Pred. No. 7.54e-01;
Matches 7; Conservative 4; Mismatches 6; Indels 0; Gaps 0;

Db 450 idhlgrrrvrcvgemae 466

||||| : :||| :|

Qy 2 IDHGTGKSKVRQKVE 18

RESULT 10

ID YTX1 XENLA STANDARD; PRT; 775 AA.

AC P14380;

DT 01-JAN-1990 (REL. 13, CREATED)

DT 01-JAN-1990 (REL. 13, LAST SEQUENCE UPDATE)

DT 01-APR-1990 (REL. 14, LAST ANNOTATION UPDATE)

DE TRANSPOSON TX1 HYPOTHETICAL 82 KD PROTEIN (ORF 1).

OS XENOPUS LAEVIS (AFRICAN CLAWED FROG).

OC EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; AMPHIBIA; ANURA.

RN [1]

RP SEQUENCE FROM N.A.

RX MEDLINE; 89384562.

RA GARRETT J.E., KNUZZON D.S., CARROLL D.;

RL MOL. CELL. BIOL. 9:3018-3027(1989).

DR EMBL; M26915; G214845; -.

DR PIR; A32494; A32494.

KW HYPOTHETICAL PROTEIN; TRANSPOSABLE ELEMENT.

SQ SEQUENCE 775 AA; 82355 MW; 9738B05A CRC32;

Query Match 39.0%; Score 55; DB 11; Length 775;

Best Local Similarity 46.7%; Pred. No. 1.24e+00;

Matches 7; Conservative 5; Mismatches 3; Indels 0; Gaps 0;

Db 617 sntskcvsevegt 631

||||| |||:

Qy 7 TKSSKVRQKVEGSS 21

RESULT 11

ID COTZ B-CSI STANDARD; PRT; 148 AA.

AC Q08312;

DT 01-OCT-1994 (REL. 30, CREATED)

DT 01-OCT-1994 (REL. 30, LAST SEQUENCE UPDATE)

DT 01-FEB-1995 (REL. 31, LAST ANNOTATION UPDATE)

DE SPORE COAT PROTEIN Z.

GN COTZ.

OS BACILLUS SUBTILIS.

OC PROKARYOTA; FIRMICUTES; ENDOSPORE-FORMING RODS AND COCCI; BACILLACEAE.

RN [1]

RP SEQUENCE FROM N.A.

RC STRAIN=168 / JH642;

RX MEDLINE; 93285989.

RA ZHANG J., FITZ-JAMES P.C., ARONSON A.I.;

RL J. BACTERIOL. 175:3757-3766(1993).

CC -!- SUBCELLULAR LOCATION: SPORE OUTER COAT.

CC -!- SUBUNIT: DISULFIDE CROSS-LINKED EITHER TO ITSELF OR TO COTY.

CC -!- SIMILARITY: TO COTY.

DR EMBL; L10116; G304149; -.

DR PIR; E47119; E47119.

DR SUBTILIST; BG10500; COTZ.

KW SPORULATION.

SQ SEQUENCE 148 AA; 16534 MW; B5442F5E CRC32;

Query Match 38.3%; Score 54; DB 2; Length 148;

Best Local Similarity 63.6%; Pred. No. 2.01e+00;

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13

RX MEDLINE; 83164198.
RA SARNOW P., HEARING P., ANDERSON C.W., REICH N., LEVINE A.J.;
RL J. MOL. BIOL. 162:565-583(1982).
RN [2]
RP COMPLETE GENOME.
RX MEDLINE; 92087470.
RA CHROBOCZEK J., BIEBER F., JACROT B.;
RL VIROLOGY 186:280-285(1992).
DR EMBL; M73260; -: NOT_ANNOTATED_CDS.
DR EMBL; X02998; G58502; -.
DR PIR; B03807; Q4ADE5.
KW EARLY PROTEIN.
SQ SEQUENCE 116 AA; 13298 MW; 66EA9B5C CRC32;

Query Match 37.6%; Score 53; DB 3; Length 116;
Best Local Similarity 60.0%; Pred. No. 3.25e+00;
Matches 6; Conservative 3; Mismatches 1; Indels 0; Gaps 0;

Db 3 rclrlkvega 12
:|:| | | | |
Qy 11 KCVQRKVEGS 20

RESULT 15
ID E411 ADE02 STANDARD; PRT; 116 AA.
AC P03241;
DT 21-JUL-1986 (REL. 01, CREATED)
DT 21-JUL-1986 (REL. 01, LAST SEQUENCE UPDATE)
DT 01-APR-1990 (REL. 14, LAST ANNOTATION UPDATE)
DE PROBABLE EARLY E4 11 KD PROTEIN.
OS HUMAN ADENOVIRUS TYPE 2.
OC VIRIDAE; DS-DNA NONENVELOPED VIRUSES; ADENOVIRIDAE; MASTADENOVIRUSES.
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE; 82059444.
RA HERRISSE J., RIGOLET M., DUPONT DE DINECHIN S., GALIBERT F.;
RL NUCLEIC ACIDS RES. 9:4023-4042(1981).
DR EMBL; J01917; G209839; -.
DR PIR; A03807; Q4ADE2.
KW EARLY PROTEIN.
SQ SEQUENCE 116 AA; 13255 MW; 950D6981 CRC32;

Query Match 37.6%; Score 53; DB 3; Length 116;
Best Local Similarity 60.0%; Pred. No. 3.25e+00;
Matches 6; Conservative 3; Mismatches 1; Indels 0; Gaps 0;

Db 3 rclrlkvega 12
:|:| | | | |
Qy 11 KCVQRKVEGS 20

Search completed: Wed Jan 28 12:11:02 1998
Job time : 10 secs.

Result	No.	Score	Query Match	Length	DB	ID	Description	Pred. No.
1	141	100.0	21	15	R77605		Pro-C5 polypeptide KS	1.23e-09
2	141	100.0	1676	15	R77604		Pro-C5 polypeptide.	1.23e-09
3	55	39.0	264	4	R2271		CSHase.	3.50e+01
4	54	38.3	1732	17	R96029		P. gingivalis porphyr	4.47e+01
5	54	38.3	3163	16	R94347		Hepatitis GB virus (H	4.47e+01
6	53	37.6	559	3	R14327		Mouse epithelin precu	5.69e+01
7	53	37.6	652	15	R88124		Tobacco mosaic virus	5.69e+01
8	53	37.6	1009	5	R26706		Type B human platelet	5.69e+01
9	53	37.6	1089	2	R06910		Alpha type PDGF recep	5.69e+01
10	53	37.6	1089	2	R08267		Platelet derived grow	5.69e+01
11	53	37.6	1196	19	W04326		Rat petrin.	5.69e+01

RESULT	1
ID	R77605 standard; Protein; 21 AA.
AC	R77605;
DT	02-APR-1996 (first entry)
DE	Pro-C5 polypeptide KSSKC epitope.
KW	Complement C5; haemolysis; kidney; glomerulonephritis;
KW	monoclonal antibody; antiinflammatory; antibody engineering;
KW	humanised antibody; KSSKC epitope.
OOS	Homo sapiens.
PEN	W09529697-A1.
PPD	09-NOV-1995.
PPF	01-MAY-1995; U05688.
PPR	02-MAY-1994; US-236208.
PPA	(ALEX-) ALEXION PHARM INC.
PPI	Evans MJ, Matis L, Moeller EE, Nye SH, Rollins S;
PPI	Rother RP, Springhorn J P, Squinto SP, Thomas TC;
PPI	Wang Y, Wilkins JA;
DR	WPI; 95-392923/50.
PPT	Treating glomerulonephritis with antibody against complement C5
PPT	component - to inhibit complement induced cell lysis
PPT	Example 13; Page 81; 181pp; English.
CCC	The cDNA sequence of the complement C5 gene transcript predicts a
CCC	secreted pro-C5 precursor of 1676 amino acids (R77604). C5 is a
CCC	beta-globulin heterodimer thought to play a role in the pathogenesis
CCC	of glomerulonephritis (GN). Cleavage of the C5 alpha-chain
CCC	by a convertase enzyme generates anaphylatoxic C5a. Monoclonal

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3

CC and humanised recombinant antibodies that recognise the alpha-chain
 CC KSKC epitope (R77605) block C5a generation, thereby reducing
 CC glomerular inflammation and kidney dysfunction associated with GN.
 SQ Sequence 21 AA;

Query Match 100.0%; Score 141; DB 15; Length 21;
 Best Local Similarity 100.0%; Pred. No. 1.23e-09;
 Matches 21; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Db 1 vidhgktkaskcvrkvegss 21
 |||||
 Qy 1 VIDHGTGKSKCVRKVEGSS 21

RESULT 2

ID R77604 standard; Protein; 1676 AA.
 AC R77604;
 DT 15-MAR-1996 (first entry)
 DE Pro-C5 polypeptide.
 KW Complement C5; haemolysis; kidney; glomerulonephritis;
 KW monoclonal antibody; antiinflammatory; antibody engineering;
 KW humanised antibody.
 OS Homo sapiens.
 FH Key Location/Qualifiers
 FT Peptide 1..18
 FT /label= Sig_peptide
 FT Protein 19..673
 FT /label= Beta-chain
 FT Cleavage_site 673..674
 FT Cleavage_site 677..678
 FT Peptide 674..677
 FT label= Cleavage_peptide
 FT Protein 678..1676
 FT /label= Alpha-chain
 FT /note= "amino acids 872-892 (854-874 of
 FT the mature protein) comprise the KSKS
 FT epitope".
 FT Peptide 678..751
 FT /label= C5a
 FT Cleavage_site 751..752
 FT /label= Convertase_cleavage_site
 FT Modified_site 911
 FT /label= N-glycosylation_site
 FT Modified_site 1115
 FT /label= N-glycosylation_site
 FT Modified_site 1630
 FT /label= N-glycosylation_site
 FT W09529697-Al.
 PD 09-NOV-1995.
 PF 01-MAY-1995; U05688.
 PR 02-MAY-1994; US-236208.
 PA (ALEX-) ALEXION PHARM INC.
 PI Evans MJ, Matlis L, Mueller EE, Nye SH, Rollins S;
 PI Rother RP, Springhorn J P, Squinto SP, Thomas TC;
 PI Wang Y, Wilkins JA;
 DR WPI; 95-392923/50.
 PT Treating glomerulonephritis with antibody against complement C5
 PT component - to inhibit complement induced cell lysis
 PS Example 13; Page 82-92; 181pp; English.
 CC The cDNA sequence of the complement C5 gene transcript predicts a
 CC secreted pro-C5 precursor of 1676 amino acids (R77604). C5 is a
 CC beta-globulin heterodimer thought to play a role in the pathogenesis
 CC of glomerulonephritis (GN). Cleavage of the C5 alpha-chain
 CC by a convertase enzyme generates anaphylatoxic C5a. Monoclonal

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US-08-487-283A-1.fag

4

CC and humanised recombinant antibodies that recognise the alpha-chain
 CC KSKC epitope (R77605) block C5a generation, thereby reducing
 CC glomerular inflammation and kidney dysfunction associated with GN.
 SQ Sequence 1676 AA;

Query Match 100.0%; Score 141; DB 15; Length 1676;
 Best Local Similarity 100.0%; Pred. No. 1.23e-09;
 Matches 21; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Db 872 vidhgktkaskcvrkvegss 892
 |||||
 Qy 1 VIDHGTGKSKCVRKVEGSS 21

RESULT 3

ID R22271 standard; Protein; 264 AA.
 AC R22271;
 DT 30-JUL-1992 (first entry)
 DE CSHase.
 KW N-carbamoyl-sarcosine amidohydrolase; CSH; assay; diagnosis;
 KW creatinine.
 OS Arthrobacter sp. DSM 2563.
 PN EP-476670-A.
 PD 25-MAR-1992.
 PF 19-SEP-1991; 115974.
 PR 20-SEP-1990; DE-029844.
 PA (BOEF) BOEHRINGER MANNHEIM GMBH.
 PI Bertscher H, Schumacher G;
 DR WPI; 92-098378/13.
 DR N-PSDB; Q22713.
 PT Recombinant DNA encoding N-carbamoyl-sarcosine-amidohydrolase -
 PT useful in clinical assay of creatinine, and vectors providing
 PT efficient expression in E.coli
 PS Claim 9; Page 9 + 7; 12pp; German.
 CC The sequence encoding CSHase is useful in assay of creatinine
 CC (for diagnosis of kidney disease). It can now be prepd. more
 CC simply than by known methods which involve culture of Arthrobacter
 CC on complex media.
 SQ Sequence 264 AA;

Query Match 39.0%; Score 55; DB 4; Length 264;
 Best Local Similarity 46.2%; Pred. No. 3.50e+01;
 Matches 6; Conservative 4; Mismatches 3; Indels 0; Gaps 0;

Db 171 gataagcvrhtve 183
 | : : ||| : ||
 Qy 6 GTKSKCVRKVE 18

RESULT 4

ID R96029 standard; Protein; 1732 AA.
 AC R96029;
 DT 04-SEP-1996 (first entry)
 DE P. gingivalis porphyain.
 KW Porphyain; haemagglutinin; periodontal disease; vaccine; antibody.
 OS Porphyromonas gingivalis strain W12.
 FH Key Location/Qualifiers
 FT Region 688..708
 FT /note= "Pro-Asn repeat region type 1"
 FT Region 887..952
 FT /note= "Pro-Asn repeat region type 2"
 FT Region 946..967
 FT /note= "Pro-Asn repeat region type 1"
 FT Region 985..1006

17-AUG-1995.
14-FEB-1995; U02118.
14-FEB-1994; US-196030.
13-MAY-1994; US-242654.
29-JUL-1994; US-283314.

Key	Location/Qualifiers
Protein	1..589
/label= "precursor"	
/note= "claim 21, page 55"	
Protein	280..335
/label= EP-1	
/note= "claim 22, page 55"	
Protein	205..261
/label= EP-2	
/note= "claim 23, page 55"	
Peptide	59..114
/label= EP	
/note= "claim 24, page 55"	
Peptide	123..179
/label= EP	
/note= "claim 25, page 55"	
Peptide	362..416
/label= EP	
/note= "claim 26, page 56"	
Peptide	440..495
/label= EP	
/note= "claim 27, page 56"	
Peptide	515..570
/label= EP	

FT /note= "claim 28, page 56"
 PN WO9115510-A.
 PN 17-OCT-1991.
 PD PF 03-APR-1991; U02321.
 PD PF 03-APR-1990; US-504508.
 PR PR 13-MAR-1991; US-083796.
 PA (BRIM) BRISTOL-MYERS SQUIB.
 PI Shoyab M, Plowman GD;
 DR WPI; 91-325168/44.
 DR N-PSDB; Q14340.
 PT New cysteine-rich growth modulating proteins, epithelins - useful
 PT as inhibitors of neoplastic cell growth and to promote wound
 PT healing and treat psoriasis
 PS Disclosure; Fig 23; 97pp; English.
 CC ET-1 is a bifunctional growth regulator, capable of stimulating
 CC the growth of some cell types while inhibiting the growth of others.
 CC ET-2 is functionally similar to ET-1 w.r.t. growth inhibitory
 CC bioactivity. In contrast, however, ET-2 is apparently not capable of
 CC eliciting the growth stimulatory activity characteristic of ET-1 and,
 CC in fact, antagonises this ET-1 activity.
 CC See also Q14338-40, Q14952-53, R14328-9 and R15315-20.
 SO Sequence 589 AA;

Query Match 37.6%; Score 53; DB 3; Length 589;
Best Local Similarity 35.7%; Pred. No. 5.69e+01;
Matches 5; Conservative 7; Mismatches 2; Indels 0; Gaps 0;

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Db 560 hcsargtkclrki 573
    | : : : : | : | :
Ov 4 HQGTSKSCVRQKV 17

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RESULT	7	
ID	R88124	standard; Protein; 652 AA.
AC	R88124;	
DT	28-MAR-1996	(first entry)
DE	Tobacco mosaic virus resistance N gene truncated protein.	
KW	Tobacco mosaic virus resistance; TMV; N gene; Solanaceae;	
KW	crop improvement; transgenic plant; crop improvement.	
OS	Nicotiana glutinosa.	
PN	MO935024-A1.	
PD	28-DEC-1995.	
PF	16-JUN-1995; U07754.	
PR	17-JUN-1994; US-261663.	
PA	(REGC) UNIV CALIFORNIA.	
PA	(USDA) US SEC OF AGRIC.	
PI	Baker BJ Whitham SA;	
DR	MPI; 96-058144/06.	
DR	N-PSDB; TQ9342.	
PT	Plant virus resistance gene N sequences from Tobacco - useful for	
PT	generating transgenic Solanaceous plants resistant to Tobacco Mosaic	
PT	Virus	
PS	Claim 28; Page 75-79; 98pp; English.	
CC	The Nicotiana glutinosa N gene truncated protein (R88124) mediates	
CC	resistance to tobacco mosaic virus (TMV). A cDNA clone (TQ9342)	
CC	coding for the protein was obtd. from a N. glutinosa leaf cDNA	
CC	library by transposon tagging. DNA sequences encoding the	
CC	protein can be used to generate transgenic plants, esp. Solanaceae,	
CC	resistant to TMV.	
SQ	Sequence 652 AA;	

Query Match 37.6%; Score 53; DB 15; Length 652;
Best Local Similarity 31.3%; Pred. No. 5.69e+01;
Matches 5; Conservative 5; Mismatches 6; Indels 0; Gaps 0;

Db 156 dnrdktddacirqivd 171
|:: : l:l l:
Qy 3 DHQGTSSKRCVRQKE 18

RESULT 8
ID R26206 standard; Protein: 1009 AA.

RESULT	8
ID	R26206 standard; Protein; 1009 AA.
AC	R26206;
AD	
DT	09-FEB-1993 (first entry)
DE	Type B human platelet-derived growth factor receptor.
KW	PDGF; PDGF-R; mesenchyme; tyrosine kinase; ligand binding region.
KE	
OS	Homo sapiens.
FT	Key
FT	Location/Qualifiers
FT	Peptide
FT	1..23
FT	/label= Signal_peptide
FT	Protein
FT	24..1009
FT	/label= Mature_PDGF-A
PN	W09213867-A.
PD	20-AUG-1992.
PF	28-JAN-1992; U00730.
PR	31-JAN-1991; US-650793.
PT	(CORT-) COR THERAPEUTICS INC.
PA	Escobedo JA, Fretto LJ, Giese NA, Tomlinson JE, Williams LT;
PI	Wolf D;
PI	WPI; 92-299970/36.
DR	N-PSDB; Q27451.
DR	
PT	Platelet derived growth factor receptor (PDGF-R) poly:peptide(s)
PT	- useful as therapeutic and diagnostic agents e.g. for assaying
PT	PDGF activity in sample
PS	Disclosure; Page 90; 109pp; English.
CC	The sequence given is one allele of type A human platelet-derived
CC	growth factor (PDGF) receptor (PDGF-R). This receptor is typical
CC	found on cells of mesenchymal origin. It acts while in the form of
CC	two transmembrane glycoproteins, each of which is about 180 kD.
CC	This receptor has three major regions. The first is a transmembrane
CC	region, which spans the membrane once, separating the regions of the
CC	receptor exterior to the cell from those interior to the cell. The
CC	second region is an extracellular region which contains the domain
CC	which bind the PDGF. The third region is an intracellular region
CC	which possesses a tyrosine kinase activity. This tyrosine kinase
CC	domain is notable in having an insert of approx. 100 amino acids,
CC	as compared with most other receptor tyrosine kinase domains which
CC	are contiguous or have shorter insert sequences. Fragments of this
CC	sequence between 8 and 400 amino acids comprising one or more PDGF
CC	ligand binding region from the extracellular domain may be used to
CC	bind a PDGF ligand.
CC	
SQ	Sequence 1009 AA;

Query Match	37.6%	Score 53;	DB 5;	Length 1009;
Best Local Similarity	38.1%;	Pred. No. 5.69e+01;		
Matches	8:	Conservative	6:	Mismatches
			7:	Indels
			0:	Gaps
			0:	

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Db 381 vddhgstggqtvrctaegt 401
    | | | | : : | | | | : :
Qv 1 VIDHQTSSKCVRKVEGSS 21

```

RESULT 9
ID R06910 standard; protein; 1089 AA.

DT 16-JAN-1991 (first entry)
DE Alpha type PDGF receptor deduced from TR4 cDNA clone.
KW Platelet derived growth factor; T11.

OS Homo sapiens. Location/Qualifiers
 FH Key 1..23
 FT Domain
 FT /label=signal peptide
 FT Domain 24..524
 FT /label=ligand binding domain
 FT Domain 525..548
 FT /label=transmembrane region
 FT Domain 549..599
 FT /label=juxtamembrane domain
 FT Binding-site 600..627
 FT /label=ATP binding site
 FT Modified-site 849
 FT /label=tyrosine autophosphorylation site
 FT Modified-site 42..44
 FT /label=N-glycos_site
 FT Modified-site 76..78
 FT /label=N-glycos_site
 FT Modified-site 103..105
 FT /label=N-glycos_site
 FT Modified-site 179..181
 FT /label=N-glycos_site
 FT Modified-site 353..355
 FT /label=N-glycos_site
 FT Modified-site 359..361
 FT /label=N-glycos_site
 FT Modified-site 458..460
 FT /label=N-glycos_site
 FT Modified-site 468..470
 FT /label=N-glycos_site
 PN W09010013-A.
 PD 07-SEP-1990.
 PF 08-FEB-1990; U00617.
 PR 09-FEB-1989; US-308282.
 PA (USDC) US SEC OF COMMERCE.
 PI Matsui T, Aaronson SA, Pierce JH;
 DR WPI; 90-290306/38.
 DR N-PSDB; Q05989.
 PT Type alpha platelet-derived growth factor :ceptor gene - useful
 PT for transforming cells to express novel protein receptor and also
 PT susceptible to genetic engineering.
 PS Claim 7; Fig 3; 64pp; English.
 CC The TR4 clone is the largest cDNA clone related to the T11 genomic
 CC clone. Isolated from a library prepd. from human thymus DNA. The
 CC T4 cDNA clone was isolated from a M426 human embryo fibroblast
 CC cDNA library. The coding region can be introduced into the pSV2
 CC gpt vector with a simian sarcoma virus LTR as a promoter and
 CC expressed in a host. The resulting protein is a novel PDGF
 CC receptor designated type alpha (the known receptor is designated
 CC type beta). The polypeptide has a calculated molecular mass of 120
 CC kD and has all the characteristics of a membrane spanning tyrosine
 CC kinase receptor. The extracellular region comprises a hydrophobic
 CC signal peptide and a ligand binding domain which has structural
 CC homology with the PDGF-R/CSF1-R subfamily. Ten Cys residues are
 CC spaced at the same positions as in other receptors of the sub-
 CC family and eight potential N-linked glycosylation sites are also
 CC present. A hydrophobic segment spans the membrane and the cyto-
 CC plasmic region comprises a juxtamembrane region, a tyrosine kinase
 CC region split into TK1 and TK2 by a hydrophilic interkinase region
 CC and a hydrophilic C-terminal tail. The TK region includes the
 CC consensus ATP binding sequence (G-X-G-X-G...K) and a tyrosine
 CC autophosphorylation site homologous to that of pp60(v-src).
 SQ Sequence 1089 AA;

Query Match 37.6%; Score 53; DB 2; Length 1089;
 Best Local Similarity 38.1%; Pred. No. 5.69e+01;
 Matches 8; Conservative 6; Mismatches 7; Indels 0; Gaps 0;
 Db 421 vddhgstggtqvtvrtaeqtp 441
 I |||: : : || ||: :
 Qy 1 VIDHQGTSKSKVQRKVEGSS 21
 RESULT 10
 ID R08267 standard; protein; 1089 AA.
 AC R08267;
 DT 07-MAR-1991 (first entry)
 DE Platelet derived growth factor (PDGF) receptor protein.
 KW Atherosclerosis; fibrotic diseases.
 OS Homo sapiens.
 PN W09014425-A.
 PD 29-NOV-1990.
 PF 21-MAY-1990; U02849.
 PR 22-MAY-1989; US-355018.
 PA (ZYMO-) ZYMOGENETICS INC.
 PI Kelly JD, Murray MJ;
 DR WPI; 90-375992/50.
 DR N-PSDB; Q06869.
 PT DNA encoding platelet-derived growth factor - used to transform
 PT cells for culturing to detect PDG agonists and antagonists
 PS Claim 1; Fig 1; 30pp; English.
 CC Gene product may be expressed from a transformed cell. It has
 CC utility in dection of PDGF agonist and antagonist analogues, binding
 CC AA, AB and BB isoforms. PDGF agonists may be used to enhance wound
 CC healing, and antagonists may be used to block the effects of PDGF
 CC eg. in treatment of atherosclerosis or fibrotic diseases.
 SQ Sequence 1089 AA;
 Query Match 37.6%; Score 53; DB 2; Length 1089;
 Best Local Similarity 38.1%; Pred. No. 5.69e+01;
 Matches 8; Conservative 6; Mismatches 7; Indels 0; Gaps 0;
 Db 421 vddhgstggtqvtvrtaeqtp 441
 I |||: : : || ||: :
 Qy 1 VIDHQGTSKSKVQRKVEGSS 21
 RESULT 11
 ID W04326 standard; Protein; 1196 AA.
 AC W04326;
 DT 16-JAN-1997 (first entry)
 DE Rat petrin.
 KW Petrin; neurite outgrowth associated protein; CNS;
 KW central nervous system; myelin; protein phosphatase 2C; stroke;
 KW neurodegeneration.
 OS Rattus sp.
 FH Key Location/Qualifiers
 FT Misc_difference 129
 FT /note= "corresponds to stop codon in DNA sequence"
 FT Misc_difference 192
 FT /note= "corresponds to stop codon in DNA sequence"
 FT Misc_difference 205
 FT /note= "corresponds to stop codon in DNA sequence"
 FT Misc_difference 219
 FT /note= "corresponds to stop codon in DNA sequence"
 FT Misc_difference 225
 FT /note= "corresponds to stop codon in DNA sequence"
 FT Misc_difference 234

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US-08-487-283A-1,100

Qy 11 KQVQRKVESS 21

RESULT 13
 ID R14325 standard; Protein; 589 AA.
 AC R14325;
 DT 17-JAN-1992 (first entry)
 DE Rat epithelin precursor.
 KW ET; growth regulation; inhibition; stimulation.
 OS Rattus rattus.
 FH Key Location/Qualifiers
 FT Protein 1..589
 FT /label= precursor
 FT /note= "claim 11, page 54"
 FT Protein 280..335
 FT /label= EP-1
 FT /note= "claim 12, page 54"
 FT Protein 205..261
 FT /label= EP-2
 FT /note= "claim 13, page 54"
 FT Peptide 59..114
 FT /label= EP
 FT /note= "claim 14, page 54"
 FT Peptide 123..179
 FT /label= EP
 FT /note= "claim 15, page 54"
 FT Peptide 362..416
 FT /label= EP
 FT /note= "claim 16, page 54"
 FT Peptide 440..495
 FT /label= EP
 FT /note= "claim 17, page 54"
 FT Peptide 515..570
 FT /label= EP
 FT /note= "claim 18, page 55"
 PN W09115510-A.
 PD 17-OCT-1991.
 PF 03-APR-1991; U02321.
 PR 03-APR-1990; US-504508.
 PR 13-MAR-1991; US-083796.
 PA (BRIM) BRISTOL-MYERS SQUIB.
 PI Shoyab M, Plowman GD;
 DR WPI; 91-325168/44.
 DR N-PSDB; Q14338.
 PT New cysteine-rich growth modulating proteins, epithelins - useful
 PT as inhibitors of neoplastic cell growth and to promote wound
 PT healing and treat psoriasis
 PS Disclosure; Fig 18; 97pp; English.
 CC ET-1 is a bifunctional growth regulator, capable of stimulating
 CC the growth of some cell types while inhibiting the growth of others.
 CC ET-2 is functionally similar to ET-1 w.r.t. growth inhibitory
 CC bioactivity. In contrast, however, ET-2 is apparently not capable of
 CC eliciting the growth stimulatory activity characteristic of ET-1 and,
 CC in fact, antagonises this ET-1 activity.
 CC See also Q14338-40, Q14952-53, R14328-9 and R15315-20.
 SQ Sequence 589 AA;

Query Match 36.2%; Score 51; DB 3; Length 589;
 Best Local Similarity 46.2%; Pred. No. 9.21e+01;
 Matches 6; Conservative 5; Mismatches 2; Indels 0; Gaps 0;
 Db 560 hcaagtkclrrk 572
 | ::::|:| |
 Qy 4 HQGTRSKVRQK 16

RESULT 14
 ID R15785 standard; Protein; 914 AA.
 AC R15785;
 DT 10-FEB-1992 (first entry)
 DE B.thuringiensis toxin/AcNPV gp64 fusion protein.
 KW chimeric; fusion protein; insecticide; AcNPV; Lepidoptera larvae;
 KW midgut targeting; bacterial endotoxin; pFAC13.
 OS Bacillus thuringiensis var. tenebriosis.
 OS Autographa californica Nuclear Polyhedrosis Virus.
 PN W09117254-A.
 PD 14-NOV-1991.
 PF 02-MAY-1991; U03008.
 PR 03-MAY-1990; US-518575.
 PA (REGC) UNIV OF CALIFORNIA.
 PI Sivasubramanian N, Federici A;
 DR WPI; 91-353775/48.
 DR N-PSDB; Q14808.
 PT Extending host range or toxicity of insecticidal proteins - using
 PT protein capable of binding to gut epithelium of insects
 PS Claim 55; Fig 18; 61pp; English.
 CC A polylinker was inserted into the XmnI restriction site at the
 CC carboxyl terminus coding region of B.thuringiensis var. tenebriosis
 CC (Btt) toxin. DNA encoding the gp64 viral membrane protein of AcNPV
 CC was operably linked to the Btt toxin coding sequence via the
 CC polylinker. The gp64 gene sequences act as midgut targeting
 CC signals for bacterial endotoxins. Plasmid pFAC13 was one of three
 CC different Btt/gp64 gene fusions that were constructed and its
 CC deduced amino acid sequence is given here.
 CC See also Q14806 and Q14807.
 SQ Sequence 914 AA;

Query Match 36.2%; Score 51; DB 3; Length 914;
 Best Local Similarity 45.5%; Pred. No. 9.21e+01;
 Matches 5; Conservative 5; Mismatches 1; Indels 0; Gaps 0;
 Db 665 kfnrcikrtve 675
 | ::::|:| |
 Qy 8 KSKKVRQKVE 18

RESULT 15
 ID R15784 standard; Protein; 956 AA.
 AC R15784;
 DT 10-FEB-1992 (first entry)
 DE B.thuringiensis toxin/AcNPV gp64 fusion protein.
 KW chimeric; fusion protein; insecticide; AcNPV; Lepidoptera larvae;
 KW midgut targeting; bacterial endotoxin; pF7.
 OS Bacillus thuringiensis var. tenebriosis.
 OS Autographa californica Nuclear Polyhedrosis Virus.
 PN W09117254-A.
 PD 14-NOV-1991.
 PF 02-MAY-1991; U03008.
 PR 03-MAY-1990; US-518575.
 PA (REGC) UNIV OF CALIFORNIA.
 PI Sivasubramanian N, Federici A;
 DR WPI; 91-353775/48.
 DR N-PSDB; Q14807.
 PT Extending host range or toxicity of insecticidal proteins - using
 PT protein capable of binding to gut epithelium of insects
 PS Claim 55; Fig 17; 61pp; English.
 CC A polylinker was inserted into the XmnI restriction site at the
 CC carboxyl terminus coding region of B.thuringiensis var. tenebriosis

Jan 28 12:16

US-08-487-283A-1.rag

15

CC (Btt) toxin. DNA encoding the gp64 viral membrane protein of hcnPV
CC was operably linked to the Btt toxin coding sequence via the
CC polylinker. The gp64 gene sequences act as midgut targeting
CC signals for bacterial endotoxins. Plasmid pFX7 was one of three
CC different Btt/gp64 gene fusions that were constructed and its
CC deduced amino acid sequence is given here.
CC See also Q14806 and Q14808.
SQ Sequence 956 AA;

Query Match 36.2%; Score 51; DB 3; Length 956;
Best Local Similarity 45.3%; Pred. No. 9.21e+01;
Matches 5; Conservative 5; Mismatches 1; Indels 0; Gaps 0;

Db 707 kfncikrkve 717

|::|::|

Qy 8 KSKCVQKVE 18

Search completed: Wed Jan 28 12:09:58 1998
Job time : 14 secs.


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#cross-references MUID:88209511
#accession A27689
##molecule_type mRNA
##residues 412-1676 ##label WET
##cross-references GB:M18879
REFERENCE A01267
#authors Fernandez, H.N.; Hugli, T.E.
#journal J. Biol. Chem. (1978) 253:6955-6164
#title Primary structural analysis of the polypeptide portion of
human C5a anaphylatoxin. Polypeptide sequence determination
and assignment of the oligosaccharide attachment site in
C5a.
#cross-references MUID:79005687
#accession A01267
##molecule_type protein
##residues 678-751 ##label FER
REFERENCE A01266
#authors Lundwall, A.B.; Wetzel, R.A.; Kristensen, T.; Whitehead,
A.S.; Woods, D.E.; Onden, R.C.; Colten, H.R.; Tack, B.F.
#journal J. Biol. Chem. (1985) 260:2108-2112
#title Isolation and sequence analysis of a cDNA clone encoding the
fifth complement component.
#cross-references MUID:85130937
#accession A01266
##molecule_type mRNA
##residues 412-854,
'SIALSPRLCNCNKISGCHKLRIPGSSDSPASASQVAGITGTHHQAQPT'
##cross-references GB:K02874
##label LUN
##note the carboxyl-terminal part of the sequence in this
report appears to be derived from translation of an
A10 repeat sequence
REFERENCE S15121
#authors Bohnsack, J.F.; Mollison, K.W.; Buko, A.M.; Ashworth, J.C.;
Hill, H.R.
#journal Biochem. J. (1991) 273:635-640
#title Group B streptococci inactivate complement component C5a by
enzymic cleavage at the C-terminus.
#cross-references MUID:91144547
#contents annotation
#COMMENT Complement C5 contains two disulfide-linked chains, formed by
removal of four basic residues. C5 convertase releases C5a
anaphylatoxin from the amino end of the alpha chain, generating
C5b (beta and alpha' chains).
#COMMENT Activation of C5 initiates the spontaneous assembly of the late
complement component, C5-C9, into the membrane attack complex.
C5b has a transient binding site for C6. The C5b-C6 complex is
the foundation upon which the membrane attack complex is
assembled.
#COMMENT C5a has potent spasmogenic and chemotactic activity.
GENETICS
#gene GDB:C5
#map_position 9q33-9q33
#CLASSIFICATION #superfamily alpha-2-macroglobulin
#KEYWORDS complement alternate pathway; complement pathway; cytolysis;
glycoprotein; inflammation; membrane attack complex; plasma
FEATURE
1-18 #domain signal sequence #status predicted #label SIG
19-673, 678-1676 #product complement C5 #status predicted #label MAT
19-673, 752-1676 #product C5b #status predicted #label C5B
19-673 #product complement C5 and C5b beta chain #status
predicted #label C5BB
678-1676 #product complement C5 alpha chain #status predicted
```

```
#label C5A
#product C5a anaphylatoxin #status experimental #label
C5T
#product C5b alpha' chain #status predicted #label C5BA\
678-751
752-1676
567-810, 634-669,
698-724, 699-731,
711-732, 866-1527;
1101-1159,
1375-1505,
1405-1474,
1520-1525,
1532-1606,
1553-1676,
1654-1657,
741
#disulfide bonds #status predicted\
#binding site carbohydrate (Asn) (covalent) #status
experimental\
#cleavage site Arg-Leu (C5 convertase) #status
experimental\
#binding_site carbohydrate (Asn) (covalent) #status
predicted
SUMMARY #length 1676 #molecular-weight 188330 #checksum 3858
Query Match 100.0%; Score 12048; DB 2; Length 1676;
Best Local Similarity 100.0%; Pred. No. 0.00e+00;
Matches 1676; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Db 1 mqlgllclflfkgwtgqetvlsapklfrvgasenivlqvyygteafdatiskayp 60
|||||
Qy 1 MGLLGLCLFLFLCKTGWGEQTVLSAPKIFRVGASENIVIQVGYGTEAFDATISKYP 60
|||||
Db 61 dkkfssysshvhlssenkfmsailltiqpklpqggnpvsyvvlewskhfskskmpit 120
|||||
Qy 61 DKKFSYSSGRVHLSENKFSQNSAILTIQPKLPQGQNPVSIVYLEWSKHSKSRMPIT 120
|||||
Db 121 ydnqflfihbtkpvytpdqgsvkrvyeinddlkpkakretvltfdogsevdmvveidhi 180
|||||
Qy 121 YDNGFLFIHDTKPYTPDQSVKRVYSINDDLKPKAKRETVLTFIDPEGSEVDMVEIDHI 180
|||||
Db 181 giisfpdfkipenrygmwtikakykedfettgtayfevkeyvlpfhfsveieynfigy 240
|||||
Qy 181 GIISFPDFKIPSNPRYGMWTIKAKYKEDFTTGTAYFEVKEYVLPFHSVSIEPEYNFIGY 240
|||||
Db 241 knfkneitikaryfynkvvtadvitfgiredlkddqkmmqtaamqntmlingiaqv 300
|||||
Qy 241 KNFKNEITIKARYFYNKVVTADVITFGIREDLKDDQKEMMQTAMQNTMLINGIAQVT 300
|||||
Db 301 fdsatvklssyysledlnnkylviavtviestgfsaeaipegikvylspklnlva 360
|||||
Qy 301 FDSATVKLSSYYSLEDLNKKYLYIAVTVIESTGFSAEAIPEGIKVYLSPIKLNLVATP 360
|||||
Db 361 lflkpgipypikvqkdsldqlvggvpvlnaqtidvngqetsdlpskvttrvddqvaf 420
|||||
Qy 361 LFLKPGIPYPINQVQKDSLDQLVGGVPVILNAQTIDVNGQETSDLPSSKVTTRVDDGVAF 420
|||||
Db 421 vlnlpsgvrtvlefnvktadpdlpeengaregyraiayslsqeylyidwtdhnhkallvge 480
|||||
Qy 421 VLNLP SGVTVLEFNKTVADPDLPEENQAREGYRAIAYSSLSQSYLYIDWTDNHKALLVGE 480
|||||
Db 481 hlniivtkspydikthylnliskgkiihfgtrekfsdasyqiniptqnmvpsarl 540
|||||
Qy 481 HLNIVTKSPYDIKITHYLNILSKGKIIHFCTREKFSDSASQSQINIPVTQNMVPSRL 540
|||||
Db 541 lvyvitytgeqtaelvdsvwlneeckcqnqlqwhlspdadavspqgtvslnmatgmdaw 600
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Qy 541 LVYYVTGQTAELVSDSWLNIIEKCGNQQLVHLSDDADAVSGQTVSLNATGDSHV 600
Db 601 alaavdsavqvqrqakklervfgfleksdlgcagagglmnanvfhlagltfltnanad 660
Qy 601 ALAAVDSAVYGVORGAKRQLERVQFLEKSDLGCCAGGLNANVFHLAGLTLTNANAD 660
Db 661 dsqendepckeilprprtqlqkkeeiaakylhsvtkccydgacvnnndetceqraariel 720
Qy 661 DSQENDEPCKEILPRPRTLQKKIEETAAKYHSHVVKCCYDCAVCNNDETCEQRAARISL 720
Db 721 gprcikafteccvvaqlranishkmdqlgrlhmklllpvskpeirsypeswlewhlv 780
Qy 721 GPRCIKAFTECCVAVASQJLANISHKMDQLGRHLHMKTLPLVSKPEIRSYPPESWLWEVHLV 780
Db 781 prtkqlqfalpdsittweiqgigientgicvadtkakvdkdfilemniyyavvrgeqig 840
Qy 781 PRRKQLQFALPDSLTTWEIQGIGISNTGICVADTKAKVDFKDFLENNIPYSVVRGEQIQ 840
Db 841 lkgtvynvrtagmqfcvkmavegiictseespwichggtksekczrqkvegsehlvtftv 900
Qy 841 LKGTVTYNVRTSGQFCVKMSAVEGICTSESPVIDHOGIKESKCVRKVEGSSHLVTFTV 900
Db 901 lpleiglhmfnsetwfgkailvktlrvpvgvkresyvgvtdlprgiygticisrrkefp 960
Qy 901 LPLEIGLHMNFNSETWFGKAILVKTLRVPEGVKRESYVGVTDLPRGIYGTISRKEFP 960
Db 961 yripdlvpkteikrlvavkglvgeilvavlsqeginlthlpkgsaeelmsavvpvfy 1020
Qy 961 YRIPDLVPKTEIKRILSVKGLVGEILSVLSQEGINLTHLPKGSAEELMSVVPVY 1020
Db 1021 vfhyletgnhwnifhsdpLIEQKQKAKKLGKGLMSIMSRNADYSYVWKGGSASTWLT 1080
Qy 1021 VFHYLETGNHWNIFHSDDLIEQKQKAKKLGKGLMSIMSRNADYSYVWKGGSASTWLT 1080
Db 1081 falrvlgqvnyveqanpsicnslilvnyqldngsfkengsqpiklqgtlpvearen 1140
Qy 1081 FALRVLGQVNYVEQANPSICNLSLILVNYQLDNGSFKENGSTQPIKLGTLPEAREN 1140
Db 1141 elyftafvigrkafdicplvkdialikadnfilentlpaqetftlaisayslgdk 1200
Qy 1141 SLVYTAFTVIGIRKAFDIPLVKIDTALIKADNFLENTLPAQSTFTLAISAYLSLGDK 1200
Db 1201 thpqfrsivskrealvkgnpvpyrfwkdlqkdsypntgtarmvettayalltsln 1260
Qy 1201 THPQFRSIVSKREALVKNPPIYRFWKDLQKDSYVPNTGTARWETTAYALLTSLN 1260
Db 1261 lkdinynvpvkwleseqrygggfyetqdtinalqgleyslvkqlrlmdidvaykhh 1320
Qy 1261 LKDINYNVPVWKWLESEQRYGGGFYSTQDTINALEGLCYSLVVKQLRLMDIDVSYKHH 1320
Db 1321 galhnykmtdknflgrpvevllnddlivstgfgslatvhvttvwhkttsetsevcfsyfk 1380
Qy 1321 GALHNYKMTDKNFLGRPVEVLLNDLLIVSTGFGSLATVHVTTVWHKTTSEEVCSFYLK 1380
Db 1381 idtqdieshyrgynsdykrivacasykpreesagshavmdislpptgisaneedlk 1440
Qy 1381 IDTQDIEASHYRGYGNSDYKRIVACASYKPRESSSSGSHAVMDISLPTGISANEEDLK 1440
Db 1441 alveqvqdlftdyqkdgdhvllqnsipsedflevfrifellevgtlspatftvyeyhr 1500
Qy 1441 ALVEGVQDLFTDQYKDGHDHVLQNSIPSEDFLEVFRIFELEVEVGLSPATFTVYEYHR 1500
Db 1501 pdkqctmftstaniqkvcegaackcveadcgmgqeeilditisaetrkqtackpeiaaya 1560
Qy 1501 PDKQCTMFTSTANIKQKVCEGAACKCVEADCGMGQEEILDITISAETRKQTACKPEIAYA 1560

Qy 1501 PKQCTMFTSTANIKQKVCEGAACKCVEADCGMGQEEILDITISAETRKQTACKPEIAYA 1560
Db 1561 ykvaiteitvenvfvykatlldiylktgeavakdseitfikkvctnaelvkgrqylim 1620
Qy 1561 YKVSITITSITVENVFVKYKATLLDIIYKKTGEAVAEKDSFITFKVCTCTNAELVKGRQYLIM 1620
Db 1621 gkealqikynfayriypdlstwieywpdrdtccsqcaflanldefaediflncg 1676
Qy 1621 GKEALQIKYNFSFRYIYPLDLSLTWIEYWPDRDTCCSSQCAFLANLDEFDAEDIFLNCG 1676
RESULT 2
ENTRY complement C5 precursor - mouse
TITLE C5a anaphylatoxin; C5b
CONTAINS
ORGANISM #formal name Mus musculus #common name house mouse
DATE 19-Nov-1988 #sequence_revision 15-Oct-1994 #text_change 16-Feb-1997
ACCESSIONS A35530; A27538; A40429
REFERENCE A35530
#authors Wetzel, R.A.; Fleischer, D.T.; Haviland, D.L.
#journal J. Biol. Chem. (1990) 265:2435-2440
#title Deficiency of the murine fifth complement component (C5). A 2-base pair gene deletion in a 5'-exon.
#cross-references MUID:90153853
#accession A35530
#molecule type mRNA
#residues 1-215,'L' #label WET
#cross-references GB:J05234
REFERENCE A27538
#authors Wetzel, R.A.; Ogata, R.T.; Tack, B.F.
#journal Biochemistry (1987) 26:737-743
#title Primary structure of the fifth component of murine complement.
#cross-references MUID:87185363
#accession A27538
#molecule type mRNA
#residues 'PCL', 44-1680 #label WET2
REFERENCE A40429
#authors Haviland, D.L.; Haviland, J.C.; Fleischer, D.T.; Wetzel, R.A.
#journal J. Biol. Chem. (1991) 266:11818-11825
#title Structure of the murine fifth complement component (C5) gene. A large, highly interrupted gene with a variant donor splice site and organizational homology with the third and fourth complement component genes.
#cross-references MUID:91268053
#accession A40429
#molecule type DNA
#residues 1-15 #label HAV
#cross-references GB:M64852
COMMENT Complement C5 contains two disulfide-linked chains, formed by removal of four basic residues. C5 convertase releases C5a anaphylatoxin from the amino end of the alpha chain, generating C5b (beta and alpha' chains).
COMMENT Activation of C5 initiates the spontaneous assembly of the late complement components, C5-C9, into the membrane attack complex. C5b has a transient binding site for C6. The C5b-C6 complex is the foundation upon which the membrane attack complex is assembled.
COMMENT C5a has potent spasmogenic and chemotactic activity.
GENETICS
#map_position 2
#introns 22/3; 86/3; 140/3; 164/3; 195/2; 223/1; 253/2; 291/3; 334/1; 372/3; 434/3; 502/3; 572/3; 622/3; 667/1; 691/1; 757/1;


```
Qy 1377 FYIKIDTDIEAS-HYRGYNSDYKRVACASYRP-SREESSGSHVMDISLPTGISAN 1435
Db 1440 eedlralvegvdqlltdykqkghvilqlnsipsrdflvrfifelfqvglfnpatftv 1499
Qy 1436 EEDLALVEGVQDLFTDQYIKDGHVILQLNSIPSSDFLCVRFIRFELFEVGLSPATFTV 1495
Db 1500 yeyhrpdkmtiysiadtrlqkvegaactcveadcaqlqaevdlaiderkackp 1559
Qy 1496 YEYHRDPKQCTMFYTSNLIKIKQVCEGACKVEADCGQMQEIDLITISAEYRKOTACKP 1555
Db 1560 etayaykvritsaateenvfkytatllytkytea-adensevtfikmcsctnanlvkqk 1618
Qy 1556 EIAYARKVSTITVENVFRKATLIDYKTEGAEVXKDSSEITFIRKVTCTNAELVRGR 1615
Db 1619 gylimgkevlqikhnfsfyjypldsatsewiywptdtcpscqafvenlmnfaedflns 1678
Qy 1616 QYLIMCKEALQIKYNSFRYIYPLDSUTWIEYPRDTTCSQCAF LANLDEFAEDIFLNG 1675
Db 1679 c 1679
Qy 1676 c 1676
```

```
RESULT 3
ENTRY C3NJ #type complete
TITLE complement C3 precursor - Naja naja
CONTAINS alternative-complement-pathway C3/C5 convertase (EC 3.4.21.47) C3b subunit; C3a anaphylatoxin
ORGANISM #formal_name Naja naja
DATE 18-Jun-1993 #sequence_revision 07-Oct-1994 #text_change 16-feb-1997
ACCESSIONS A46513
REFERENCE A46513
#authors Fritzing, D.C.; Petrella, E.C.; Connolly, M.B.; Bredehorst, R.; Vogel, C.W.
#journal J. Immunol. (1992) 149:3554-3562
#title Primary structure of cobra complement component C3.
#cross-references MUID:93054528
#accession A46513
#molecule_type mRNA
#residues 1-1651 #label FRI
#cross-references NCBIPI:118403; GB:L02365
#note authors' translation shows Arg-1408 after residue 1438 and, consequently, residues 1408-1438 are displaced one codon to the left
#note sequence extracted from NCBI backbone and corrected to correspond with the translation of the nucleotide sequence
```

COMMENT Complement C3 contains two chains, formed by removal of four residues and linked by a disulfide bond. Its activation by a C3 convertase, which is the central reaction in both classical and alternative complement pathways, releases the C3a anaphylatoxin from the amino end of the alpha chain and generates C3b, which associates with the Bb fragment of complement factor B to form the alternative-complement-pathway C3/C5 convertase.

COMMENT C3a anaphylatoxin is a vasoactive peptide and a mediator of inflammation.

COMMENT C3b, with its highly reactive thiol group, binds to the surface of foreign particles and facilitates phagocytosis. It binds to complement C5 and renders it susceptible to proteolysis by the classical-complement-pathway C3/C5 convertase. The activity of C3b is regulated by proteolytic cleavage involving factors H and I. Its degradation products can also be biologically active.

COMMENT The major site of synthesis of this plasma protein is the liver.

CLASSIFICATION #superfamily alpha-2-macroglobulin

KEYWORDS acute phase; complement alternate pathway; complement pathway; glycoprotein; hydrolase; immune response; inflammation; liver; plasma; serine proteinase; thiolester bond

```
FEATURE
1-22 #domain signal sequence #status predicted #label SIG\
23-655 #product complement C3 and C3b beta chain #status predicted #label C3BB\
23-655, 660-1651 #product complement C3 #status predicted #label CC3\
23-655, 739-1651 #product complement C3b #status predicted #label C3B\
660-1651 #product complement C3 alpha chain #status predicted #label CC3A\
660-738 #product C3a anaphylatoxin #status predicted #label C3A\
739-1651 #product complement C3b alpha' chain #status predicted #label C3BA\
1412-1445 #region properdin binding\
546-807, 615-650, 683-710, 684-717, 697-718, 863-1501, 1091-1147, 1346-1477, 1377-1446, 1494-1499, 1506-1578, 1525-1649, 1625-1634 738-739 #disulfide bonds #status predicted\
#cleavage_site Arg-Ser (C3 convertase) #status predicted\
999-1002 #cross-link thiolester (Cys-Gln) #status predicted #length 1651 #molecular-weight 184925 #checksum 3526
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SUMMARY
Query Match 19.6%; Score 2367; DB 2; Length 1651;
Best Local Similarity 30.2%; Pred. No. 0.00e+00;
Matches 518; Conservative 416; Mismatches 655; Indels 127; Gaps 97;

Db 4 malylyvaalllgfpg-s-shgalyllitpavirtdeeqilveahgdtpkeldifvhdf 61
Qy 1 MGLLGILCELI--FLGKTWGEQTVISAPKIFERVCASENIVQVYCYTEAFDTSIKSY 59
Db 62 prkqkltfgrvdmnqagsmfvtpti-kvpakelnkdkqngvrvvkvtpgqvalekvvl 120
Qy 60 PDKFESYSSGHRVHLS--SENKFNQSAILLTIQPKQLPGQNPNVSVYVLEWSKHFESKRMP 118
Db 121 layqgsvfqtqkgytpgspryrvfsvdnhmrdktviveftqpegivvs-skpvn 179
Qy 119 IYDNGELFIHTDKPVYTPDQSKVRVYSINDDLKPAKRETVLITFDPEGSEVDWVEED 178
Db 180 psgsit-p-ynlpelvsfqtwkavakyspeesyayfdvreyvlpsfevrlpsdkfl 237
Qy 179 HIGLISFPDKFIPSNRPNRGMTIKARKYKEDFTTGTATYFEWKYEVLPHPFSIEPEYNI 238
Db 238 -yidgnknfhvsiarylygkv-egvafvfgvk--i-ddakkeipdsitripidgq 292
Qy 239 GYKN-FKNEFITIKARYFNKVVTEADVITFGREDLKDDQKEMQMTAMQNTMLNGIA 297
Db 293 eatlkrdt-lrs-rfdqlnqlvghtlysvvtitegsdmvvtteggihivtspqiyft 350
Qy 298 QVTDFSETAKVLSYSLIEDIANKYLYIAVTVIESTGFSSEAEIPGKYVLSPYKILNV 357
Db 351 ktpkyfkpmpelyvvtvp-d---gs-paa-hvp-v-vs-ea--ih-segtt-lsdgt 397
Qy 358 ATPLEKPGIPYPKQVKDSDLQLVGCPVILNAQTIDVNETSDLDPSKSVTRVDGV 417
```


#journal	J. Biol. Chem. (1975) 250:8293-8301
#title	Human anaphylatoxin (C3a) from the third component of complement.
#cross-references	WUID:76069169
#accession	A92187
##molecule_type	protein
##residues	672-680, 'N', 682-699, 'Q', 701-748 ##label HUG
REFERENCE	A27603
#authors	Daoudaki, M.E.; Becherer, J.D.; Lambris, J.D.
#journal	J. Immunol. (1988) 140:1577-1580
#title	A 34-amino acid peptide of the third component of complement mediates properdin binding.
#cross-references	WUID:88154452
#accession	A27603
##molecule_type	protein
##residues	1409-1563 ##label DAO
REFERENCE	A23435
#authors	Hellman, U.; Eggertsen, G.; Engstrom, A.; Sjoquist, J.
#journal	Biochem. J. (1985) 230:353-361
#title	Amino acid sequence of the trypsin-generated C3d fragment from human complement factor C3.
#cross-references	WUID:86025442
#accession	A23435
##molecule_type	protein
##residues	1002-1012, 'E', 1014-1303 ##label HEL
##note	sequence corresponding to residues 1072-1100 was not determined but was taken from de Bruijn & Fey (reference A94605)
REFERENCE	A45830
#authors	Poznansky, M.C.; Glissold, P.M.; Lachmann, P.J.
#journal	J. Immunol. (1989) 143:1254-1258
#title	The difference between human C3F and C3S results from a single amino acid change from an asparagine to an aspartate residue at position 1216 on the alpha-chain of the complement component, C3.
#accession	A45830
##status	not compared with conceptual translation
##molecule_type	DNA
##residues	1212-1215, 'N', 1217-1223 ##label P02
##note	this is the C3S allele
#accession	B45830
##status	not compared with conceptual translation
##molecule_type	DNA
##residues	1212-1223 ##label P02
REFERENCE	S27041
#authors	Dolmer, K.; Sottrup-Jensen, L.
#journal	FEBS Lett. (1993) 315:85-90
#title	Disulfide bridges in human complement component C3b.
#contents	annotation; disulfide bonds
COMMENT	The sequence shown is the C3 fast (C3F) allele, which is found mainly in Caucasian populations and is associated with increased incidence of certain diseases.
COMMENT	Complement C3 contains two chains, formed by removal of four residues and linked by a disulfide bond. Its activation by a C3 convertase, which is the central reaction in both classical and alternative complement pathways, releases the C3a anaphylatoxin from the amino end of the alpha chain and generates C3b, which associates with the Bb fragment of complement factor B to form the alternative-complement-pathway C3/C5 convertase.
COMMENT	C3a anaphylatoxin is a vasoactive peptide and a mediator of inflammation.
COMMENT	C3b, with its highly reactive thiol group, binds to the surface of foreign particles and facilitates phagocytosis. It binds to complement C5 and renders it susceptible to proteolysis by the


```
Qy 1516 IQKVEGACACKVEADCGMQEELDLTISAETKQTACKPEIAYAYKVSITSITVENFV 1575
Db 1549 nyinsiltvkmgtdepngsnrfvshkqcdalslqkqdylvwglas-dl-wvgsr 1606
Qy 1576 KYRATLLDYTKTGEAVAKDSEITFIKKVTCTNA-ELVKGROYLIMCKEALQIKYNSFR 1634
Db 1607 fyyliiskdwleapleescqdadlqplcqdftefsdnlvifgc 1650
Qy 1635 YIYLDLSLWIEYWPRTTCCSCQ-AFL-ANLDEPAEDIFLNGC 1676

RESULT 7
ENTRY C3MS #type complete
TITLE complement C3 precursor - mouse
CONTAINS alternative-complement-pathway C3/C5 convertase (EC
3.4.21.47) C3b subunit; C3a anaphylatoxin
ORGANISM #formal name Mus musculus #common name house mouse
DATE 30-Jun-1988 #sequence_revision 30-Jun-1988 #text_change
06-Sep-1996
ACCESSIONS A92459; B92459; A92460; A93938; A21898; A54561; S16369;
S16189; I49563; I49576; A01261; A05290; A29033
REFERENCE A92459
#authors Lundwall, A.; Wetzel, R.A.; Domdey, H.; Tack, B.F.; Fey, G.H.
#journal J. Biol. Chem. (1984) 259:13851-13856
#title Structure of murine complement component C3: I. Nucleotide
sequence of cloned complementary and genomic DNA coding for
the beta chain.
#cross-references M01D:85054818
#accession A92459
#molecule_type mRNA
#residues 1-724 #label LU1
#accession B92459
#molecule_type DNA
#residues 1-124 #label LU2
REFERENCE A92460
#authors Wetzel, R.A.; Lundwall, A.; Davidson, F.; Gibson, T.; Tack,
B.F.; Fey, G.H.
#journal J. Biol. Chem. (1984) 259:13857-13862
#title Structure of murine complement component C3: II. Nucleotide
sequence of cloned complementary DNA coding for the alpha
chain.
#cross-references M01D:85054819
#accession A92460
#molecule_type mRNA
#residues 671-1663 #label WET
REFERENCE A93938
#authors Domdey, H.; Wiebauer, K.; Kazmaier, M.; Muller, V.; Odink,
K.; Fey, G.
#journal Proc. Natl. Acad. Sci. U.S.A. (1982) 79:7619-7623
#title Characterization of the mRNA and cloned cDNA specifying the
third component of mouse complement.
#cross-references M01D:83117730
#contents C3a
#accession A93938
#molecule_type mRNA
#residues 671-748 #label DOM
REFERENCE A21898
#authors Sottrup-Jensen, L.; Stepanik, T.M.; Kristensen, T.; Lohblad,
P.B.; Jones, C.M.; Wierzbicki, D.M.; Magnusson, S.; Domdey,
H.; Wetzel, R.A.; Lundwall, A.; Tack, B.F.; Fey, G.H.
#journal Proc. Natl. Acad. Sci. U.S.A. (1985) 82:9-13
#title Common evolutionary origin of alpha2-macroglobulin and
complement components C3 and C4.
#cross-references M01D:85113177
```

```
#accession A21898
#molecule_type mRNA
#residues 25-1663 #label SOT
REFERENCE A54561
#authors Hamada, J.; Cavanaugh, P.G.; Miki, K.; Nicolson, G.L.
#journal Cancer Res. (1993) 53:4418-4423
#title A paracrine migration-stimulating factor for metastatic tumor
cells secreted by mouse hepatic sinusoidal endothelial
cells: identification as complement component C3b.
#accession A54561
#molecule_type protein
#residues 25-41;749-760 #label HAM
#experimental_source migration-stimulating factor purified from medium
conditioned by mouse hepatic sinusoidal
endothelial cells
REFERENCE S16189
#authors Sato, T.; Hong, M.H.; Jin, C.H.; Ishimi, Y.; Udagawa, N.;
Shinki, T.; Abe, E.; Suda, T.
#journal FEBS Lett. (1991) 285:21-24
#title The specific production of the third component of complement
by osteoblastic cells treated with 1-alpha,
25-dihydroxyvitamin D(3).
#cross-references M01D:91293304
#accession S16369
#molecule_type protein
#residues 25-31 #label SAT
#accession S16189
#status preliminary
#molecule_type protein
#residues 671-677, 'X', 679-680 #label SA2
REFERENCE I49563
#authors Fey, G.; Domdey, H.; Wiebauer, K.; Whitehead, A.S.; Odink, K.
#journal Springer Semin. Immunopathol. (1983) 6:119-147
#title Structure and expression of the C3 gene.
#cross-references M01D:84045280
#accession I49563
#status preliminary
#molecule_type mRNA
#residues 25-136, 'Q', 138-240 #label FEY
#cross-references GB:M35659; NID:g192280; CDS_PID:g192281
REFERENCE I49576
#authors Fey, G.H.; Wiebauer, K.; Domdey, H.
#journal Ann. N. Y. Acad. Sci. (1983) 421:307-312
#title Amino acid sequences of mouse complement C3 derived from
nucleotide sequences of cloned cDNA.
#cross-references M01D:84201365
#accession I49576
#status preliminary; translated from GB/EMBL/DBJ
#molecule_type mRNA
#residues 658-761 #label RES
#cross-references GB:M30302; NID:g192391; CDS_PID:g192392
COMMENT Complement C3 contains two chains, formed by removal of four
residues and linked by a disulfide bond. Its activation by a C3
convertase, which is the central reaction in both classical and
alternative complement pathways, releases the C3a anaphylatoxin
from the amino end of the alpha chain and generates C3b, which
associates with the Bb fragment of complement factor B to form
the alternative-complement-pathway C3/C5 convertase.
C3a anaphylatoxin is a vasoactive peptide and a mediator of
inflammation.
COMMENT C3b, with its highly reactive thiol group, binds to the surface of
foreign particles and facilitates phagocytosis. It binds to
complement C5 and renders it susceptible to proteolysis by the
classical-complement-pathway C3/C5 convertase. The activity of
```



```

8
RESULT      C3CP      #type complete
ENTRY      complement C3 precursor - guinea pig
TITLE      alternative-complement-pathway C3/C5 convertase (EC
CONTAINS   3.4.21.47) C3b subunit; C3a anaphylatoxin
ORGANISM   #formal name Cavia porcellus #common name guinea pig
DATE       07-Feb-1992 #sequence_revision 07-Oct-1994 #text_change
          16-Feb-1997
ACCSSIONS  A37156; S03375; A20342; D20342; C20342; A31222
REFERENCE  A37156
          Auerbach, H.S.; Burger, R.; Dodds, A.; Colten, H.R.
          J. Clin. Invest. (1990) 86:96-106
          #journal
          #title Molecular basis of complement C3 deficiency in guinea pigs
          #cross-references MUID:90307998
          #accession A37156
          #molecule_type mRNA
          #residues 1-1666 ##label AUE
          #cross-references GB:M34054
          S03375
          Gerard, N.P.; Lively, M.O.; Gerard, C.
          Protein Seq. Data Anal. (1988) 1:473-478
          #journal
          #title Amino acid sequence of guinea pig C3a anaphylatoxin.
          #cross-references MUID:89113342
          #accession S03375
          #molecule_type protein
          #residues 676-730, 'N', 732-752 ##label GER
          #experimental source complement-activated guinea pig serum

```

#note	form isolated is inactive C3a anaphylatoxin and is missing the carboxyl-terminal arginine
REFERENCE	A90479
#authors	Thomas, M.L.; Tack, B.F.
#journal	Biochemistry (1983) 22:942-947
#title	Identification and alignment of a thiol ester site in the third component of guinea pig complement.
#cross-references	MUID:83178889
#accession	A20342
#molecule_type	protein
#residues	676-687 ##label TH1
#accession	D20342
#molecule_type	protein
#residues	993-1012,1014-1017,'E',1019-1030,'Y' ##label TH2
REFERENCE	A20342
#authors	Goldberger, G.; Thomas, M.L.; Tack, B.F.; Williams, J.J;
#journal	Colten, H.R.; Abraham, G.N.
#title	J. Biol. Chem. (1981) 256:12617-12619 NH2-terminal structure and cleavage of guinea pig pro-C3, the precursor of the third complement component.
#cross-references	MUID:82075767
#accession	C20342
#molecule_type	protein
#residues	23-38 ##label GOL
COMMENT	Complement C3 contains two chains, formed by removal of four residues and linked by a disulfide bond. Its activation by a C3 convertase, which is the central reaction in both classical and alternative complement pathways, releases the C3a anaphylatoxin from the amino end of the alpha chain and generates C3b, which associates with the Bb fragment of complement factor B to form the alternative-complement-pathway C3/C5 convertase. C3a anaphylatoxin is a vasoactive peptide and a mediator of inflammation.
COMMENT	C3b, with its highly reactive thiol group, binds to the surface of foreign particles and facilitates phagocytosis. It binds to complement C5 and renders it susceptible to proteolysis by the classical-complement-pathway C3/C5 convertase. The activity of C3b is regulated by proteolytic cleavage involving factors H and I. Its degradation products can also be biologically active.
COMMENT	The major site of synthesis of this plasma protein is the liver.
CLASSIFICATION	#superfamily alpha-2-macroglobulin
KEYWORDS	acute phase; complement alternate pathway; complement pathway; glycoprotein; hydrolase; immune response; inflammation; liver; plasma; serine proteinase; thiolester bond
FEATURE	
1-22	#domain signal sequence #status predicted #label SIG\
23-671	#product complement C3 and C3b beta chain #status predicted #label C3B8\
23-671, 676-1666	#product complement C3 #status predicted #label CC3A\
23-671, 754-1666	#product complement C3b #status predicted #label C3B\
676-1666	#product complement C3 alpha chain #status predicted #label CC3A\
676-753	#product C3a anaphylatoxin #status predicted #label C3T\
754-1666	#product complement C3b alpha' chain #status predicted #label C3BA\
951-1308	#product C3dk fragment #status predicted #label CDK\
1007-1308	#product C3d fragment #status predicted #label C3D\
1429-1461	#region properdin binding
557-821, 630-666,	
698-725, 699-732,	
712-733, 878-1517,	
1106-1163,	
1363-1493,	


```

Qy 1291 INATEGLTEYSL-LVKQLRLSMDIDVSYKHKGALHNYKMT-DKRF LGRPVEVLADDLIV 1348
Db 1314 tas-gdgkatmttltfynaqlqekancvknfhlvnsvenihlna-mg-akgalmlki-ct 1369
Qy 1349 STGFCGSLATVHTVTVHL-KTSTSEVCS-FYLIKIDTQDTEASHYRGVGNSDYKRI VACA 1406
Db 1370 ry-ly-evds-tm-tiidsmaltgflpdaedltrlekgydryierymdmmaqkavai 1424
Qy 1407 SKYSPRESSSGSSHAVMDISLPTGICANEDKALVEGVQDLFTDYQI-KD-GH---VI 1461
Db 1425 iylnkshsedeclhfkllkhfvgfipqgskvysynldektkfyhpdktgllnki 1484
Qy 1462 IQLANSIPSSDILVCRFRIFELFEVGLSPATFTVYHRPDKOCTMFYSTSN-IKI-QKV 1519
Db 1485 cignvrcagetcaslnhq-e-rldvplqiekacetndvvytkllrieedqgndiyvm 1542
Qy 1520 CEGAACKCEADCGMQEELDLTISAETRKQTACKPEIAYAKVYSITSITVENVFKYA 1579
Db 1543 dylevikqgt denpraktqoyisqrkqcealnkvndodyliwgsrdsllptk-d-kieyi 1600
Qy 1580 TLLDIYKTGEAVAEKDSITFIKKVYCTNA-ELVKGQYILMG-K-EALQIKYNSFRYI 1636
Db 1601 it-kn-twierphedecqe-eef-qklcdffaq 1630
Qy 1637 YPLDSLWTWYPRDTCSCQAFIANL-DEFAE 1669

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RESULT 11
ENTRY S21045 #type fragment
TITLE complement protein homolog - inshore hagfish (fragment)
ORGANISM #formal name Eptatretus burgeri #common_name inshore hagfish
DATE 13-Jan-1995 #sequence_revision 13-Jan-1995 #text_change 13-Jan-1995
ACCESSIONS S21045
REFERENCE Ishiguro, H.; Kobayashi, K.; Suzuki, M.; Titani, K.;
Tomonaga, S.; Kurosawa, Y.
#journal EMBO J. (1992) 11:829-837
#title Isolation of a hagfish gene that encodes a complement
component.
#cross-references M01D:92192016
#accession S21045
#status preliminary
#molecule_type mRNA
#residues 1-1620 #label ISH
#cross-references EMBL:Z11595
SUMMARY #length 1620 #checksum 3983

```

```

Query Match 17.5%; Score 2106; DB 15; Length 1620;
Best Local Similarity 29.6%; Pred. No. 0.00e+00;
Matches 466; Conservative 416; Mismatches 588; Indels 104; Gaps 88;

Db 93 ipletsrgrlvaqtdqpiytpndmirifpvtqlnpliselvldmmpdgvvvdriek 152
Qy 117 WPIYDNGLFHTDKPVTTPDQSVKRVYSLNDLAKPAKRETVLTFIDPESGEVDWEE 176

Db 153 nafevexvmelrphfpaltslgdkviksvmkdkpqnfy-eg-fkveeylptfdvzit 210
Qy 177 ID-HIG-IISPPDKIPSNPRYGMWTKA--KYKEDSTTGATAYEYKEVYVLPHPVSJSIE 232

Db 211 eegpylhvy-d-kafthikamhykpv-m-gravvrygvkhqsk-----rtllstasalar 264
Qy 233 FEYNFI-GYKNFKFETIKARYFNKRVTEADYVITFGIREDIKDDKEMQOTAMQTM 291

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Db 265 feqgeamtl-rqkhile-qvpdpkllllgslveasvisdagelensiiddipivaap 322
Qy 292 LINGIAQVTFDSEATKELSYSLIEDINKYLYLIANTVIESTGGFSEAEIPGKYVLS 351
Db 323 yslkekwtvpffkpgvpyiykvvlmpdgspagvp-i-kv-afaf-d-eegnwitqkrt 378
Qy 352 YKLNIVATPLFLKPGIPYIKVQVKDSLQQLVGVFVITLNAQTIDVNOETSDLDLP SKSVT 411
Db 379 -ndngiamqntintarnekklnikvqtederleqqaeasftiasyepsqsfihln-a- 435
Qy 412 RVDGASVFMPLSGVTVLEFNKVTADPLPEENQAREGYRAIAYSSLSQSYLYIDWTD 471
Db 436 hrevksqgehivdfiksaakdhvlfhnylmismnqklnhflqegrkgdtt--evallit 493
Qy 472 NIKALLVGEHLNIIVTPKSPYIDKITHYNYLLSKGKIHFCTREXFSDSYQSINIPVT 531
Db 494 pelvpqfrlvaffilp---egelvadsliidvkschaklaldvaggkrlfepdnmfd 550
Qy 532 QNMVPSRLLIYVYVTGEQTAELVSDSWINIEKCGNQIAQVHLSPDADAYSQCVTSIN 591
Db 551 lagesdsavagvvdkaayvldkknltankvykameasdlgcsvgsqktgplvfrdagl 610
Qy 592 MATGMSWVALAAVDSAVYGVQKAKPLERVFQFLEKSDLGCGAGGINNANVFLHAGL 651
Db 611 almakeisgmddvdkpgcpgnhttrkrelvleiaiekasty-paelrkccrdaaiesplr 669
Qy 652 TFLTNANADDSQENDEPC-KEILRPRTLAQKIE-EIAAKYKUSVVKCCYDGCACVNN-D 708
Db 670 leceertkhihdegeqogetflecckhveeelliameededlgrsggedmigeqvvi 729
Qy 709 ETCERBAARI-SIGPRCIKAFTECCV-VASQLRANISHKDMQLGRILHMKT-LLPVSKPEI 765
Db 730 rghpfesfmweiliklereaengkeritkmpdsittwdlqavevsgskglcvgspleltv 789
Qy 766 RSYFPESLWELVHLVPRR--K-QLQFA--LPDSLTTWELIQIGISNT-GICVADTVKAKV 819
Db 790 fklqfllvhtpyalkqveqlrvviynmqdvkgeiqvkcqgdigt-dae--qneplk 846
Qy 820 FKDFVLEWNPYSVVRGEIQIGTGVYNYRTSCMQFCVKMSAVEGICTSESPVHDQGTK 879
Db 847 srfav-ek-nsats-f-sfnavplassdsvs-vlarvfsgdvhdavekdrlvmpgegye 901
Qy 880 SSKVRQKVEGSSHLVTEVLPLEIGLHNIHNSLETWFGKEI--LV-KTLRVVPEGVKR 936
Db 902 emsrswsqvr-rhgg-qqvivvdnetpqnvpvgtemeafllsaqgnlvaetiqltksgk 959
Qy 937 ESYSCVTLDPGCIYGTISRKPEPYRIPDLVPKTEIKRILSVKGLLVGEILSNVLSQEG 996
Db 960 isnllrptgcqegnmmytsitvm-varylnrsdgwnkm-gdpqlkkrfsdf-itsqfas 1016
Qy 997 INILTHLPKGSAAEALM-SVVPVYVFHYLETGNHWNIFHSDPLIEKOKLKKLKEGML 1055
Db 1017 qltyrkdyeyaaahltasatwltafvakvscqqlvfpvseicgsvrwlmrk-qdkd 1075
Qy 1056 IMSYRNADYSYSVWRKGGASWLTAFALRVLGQVKNRYEQNQSICNSLILHVENYQDIN 1115
Db 1076 gsfle-ek-pvvhln-mm-gqvtkgw-ltefvfiallearesc-inevegtvrvvekah 1129
Qy 1116 GSFKENSOYQPIKLAQTLIPVEARENSLYLTAFVIGIRKAFDIPCLVKID--TALI-KAD 1172
Db 1130 gylteqamngldefplaitayalslw-kv-sdgaakvt-m-h-tl-k-tsgl-q--teel 1179
Qy 1173 NFLLENTLPAQSTFTLAI SAYALS LCKDTHPQRSIVSALKREALVKNPPIYRFWKDL 1232

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COMMENT	Residues 1446 or 1449 may be the carboxyl end of the alpha chain.
COMMENT	There are at least two genes coding for C4, C4A and C4B. Each gene has many alleles.
GENETICS	
# gene	GDB:C4A
##cross-references	GDB:IL19732
#map_position	6p21.3-6p21.3
#introns	22/2; 88/3; 156/1; 179/3; 209/2; 237/1; 269/2; 304/3; 349/1; 387/3; 447/3; 508/3; 570/3; 623/3; 666/1; 691/1; 757/1; 794/2; 818/1; 864/3; 934/3; 952/1; 1052/1; 1077/2; 1129/3; 1168/3; 1226/1; 1303/3; 1359/3; 1379/3; 1411/1; 1473/2; 1503/3; 1528/3; 1563/1; 1593/1; 1626/1; 1654/1; 1698/2
CLASSIFICATION	#superfamily alpha-2-macroglobulin
KEYWORDS	acute phase; complement classical pathway; glycoprotein; hydrolase; inflammation; plasma; polymorphism; serine proteinase; thiolester bond
FEATURE	
1-19	#domain signal sequence
20-675	#product complement C4 beta chain #status predicted #label SIG\ #label BET\
20-675; 757-1446,	#product C4b #status predicted #label C4B\
1454-1744	#product complement C4 alpha chain #status predicted #label ALP\
680-1446	#product C4a anaphylatoxin #status experimental #label C4A\
680-756	#region C4b-binding protein binding\
757-845	#product C4d fragment #status experimental #label C4D\
957-1336	#product complement C4 gamma chain #status predicted #label GAM\
1454-1744	#cleavage_site Arg-Ala (complement subcomponent C1s) #status experimental\
756-757	#cross-link thiolester (Cys-Gln) #status experimental\
1010-1013	#binding_site carbohydrate (Asn) (covalent) #status experimental
1328	#length 1744 #molecular-weight 192860 #checksum 9431
SUMMARY	
Query Match	14.0%; Score 1692; DB 2; Length 1744;
Best Local Similarity	28.2%; Pred. No. 5.65e-280;
Matches	466; Conservative 419; Mismatches 620; Indels 147; Gaps 116;
Db	139 ghlfqtqdpynpgvriryvfalqdkmpet-dtitvmvshgdlrvkkvymposi 197
Qy	124 GFLFIHTDKPYTPDQSVKRVVSYLNDLKPARETVLTFID-PEGSEVMVEEDHIGI 182
Db	198 fq-ddfvpidsepgtwksarfdsqlelnstqfvykviylnfenvkitpgkpylltp 256
Qy	193 ISFPDKIPSNPRGYMTIKAKYKEDFTTGCTAYFEKVEKVLPHFSVIEPNFI-CYK 241
Db	257 ghldemqidiqarviyqkvpgv-qgvayvrfgllde--dgkktffrglesqtklvngqshis 313
Qy	242 -NFKNFEITIKARYFNKVKVTEADVYITFGIREDKDDQKEMQMOTMONTALINGIAV 300
Db	314 lskaeqdaleklm--gtdlqglrlyvaalievpggemeeaeltswfvspfeldis 372
Qy	301 FD-SE--TAVKELSYYSIEDLNKKYLIYAVTVIESTGGFSECAEIPGIRKVLSPKMLN 357
Db	373 ktkrhlyvpafllqalvrenmspsasgi.pvkva-tvaspsvpevqdiqqntd-gsqg 430
Qy	358 ATPLFKPGIPYPIKVQVKSLDGLVGGVPVILNAQTDVNOETSDLDPSKSVTRVDDGV 417
Db	431 vsipilipqtiselqlsvsaqsh-pai--arltvaapp-sg-gpfflsierpds-rprp 484

#journal Proc. Natl. Acad. Sci. U.S.A. (1984) 81:6822-6826
#title Isolation of cDNA clones specifying the fourth component of mouse complement and its isotype, sex-limited protein.
#cross-references MUID:85038607
#accession A21692
##molecule_type mRNA
##residues 651-719, 'G', 721-738, 'AI', 741-805 ##label NO2
##cross-references GB:M12970
##experimental_source strain FM
REFERENCE
A30520
#authors Taillon-Miller, P.A.; Shreffler, D.C.
#journal J. Immunol. (1988) 141:2382-2387
#title Structural basis for the C4d.1/C4d.2 serologic allotypes of murine complement component C4.
#cross-references MUID:89009745
#accession A30520
##molecule_type DNA
##residues 961-1205, 'Q', 1207-1290 ##label TAI
##cross-references GB:M23186
##experimental_source strain B10.BR
REFERENCE
A60227
#authors Ogata, R.T.; Zepf, N.E.
#journal Eur. J. Immunol. (1990) 20:1607-1610
#title C4 from C4-high and C4-low mouse strains have identical sequences in the region corresponding to the isotype-specific segment of human C4.
#cross-references MUID:90353398
#accession A60227
##molecule_type DNA
##residues 1099-1142 ##label OGA
##cross-references GB:X55493
REFERENCE
A22039
#authors Levi-Strauss, M.; Tosi, M.; Steinmetz, M.; Klein, J.; Meo, T.
#journal Proc. Natl. Acad. Sci. U.S.A. (1985) 82:1746-1750
#title Multiple duplications of complement C4 gene correlate with H-2-controlled testosterone-independent expression of its sex-limited isoform, C4-Slp.
#cross-references MUID:85166208
#accession A22039
##molecule_type mRNA
##residues 1105-1118, 'A', 1120-1189, 'T', 1191-1449 ##label LEV
##cross-references GB:K02798
##experimental_source strain B10.W7R
REFERENCE
A93753
#authors Tosi, M.; Levi-Strauss, M.; Duponchel, C.; Meo, T.
#journal Philos. Trans. R. Soc. Lond. (1984) 306:389-394
#title Sequence heterogeneity of murine complementary DNA clones related to the C4 and C4-Slp isoforms of the fourth complement component.
#accession A29059
##molecule_type mRNA
##residues 1258-1376 ##label TOS
##cross-references GB:K02798
REFERENCE
A01264
#authors Ogata, R.T.; Shreffler, D.C.; Sepich, D.S.; Lilly, S.P.
#journal Proc. Natl. Acad. Sci. U.S.A. (1983) 80:5061-5065
#title cDNA clone spanning the alpha-gamma subunit junction in the precursor of the murine fourth complement component (C4).
#cross-references MUID:83273751
#accession A01264
##molecule_type mRNA
##residues 1360-1400, 'S', 1402-1511 ##label OCG
##cross-references GB:K00019
##experimental_source strain B10.W7R

REFERENCE
A41195
#authors Ogata, R.T.; Sepich, D.S.
#journal Proc. Natl. Acad. Sci. U.S.A. (1984) 81:4908-4911
#title Genes for murine fourth complement component (C4) and sex-limited protein (Slp) identified by hybridization to C4- and Slp-specific cDNA.
#cross-references MUID:84272739
#accession B41195
##molecule_type mRNA
##residues 1360-1400, 'S', 1402-1511 ##label OCG
##cross-references GB:K00019
##experimental_source strain B10.W7R
REFERENCE
I59084
#authors Nonaka, M.; Kimura, H.; Yeul, Y.D.; Yokoyama, S.; Nakayama, K.; Takahashi, M.
#journal Proc. Natl. Acad. Sci. U.S.A. (1986) 83:7883-7887
#title Identification of the 5'-flanking regulatory region responsible for the difference in transcriptional control between mouse complement C4 and Slp genes.
#cross-references MUID:87017050
#accession I59084
##status preliminary; translated from GB/EMBL/DBJ
##molecule_type DNA
##residues 1-21 ##label RES
##cross-references GB:M14225; NID:g199291; CDS_PID:g554211
REFERENCE
I48274
#authors Hemenway, C.; Kalff, M.; Stavenhagen, J.; Waltheil, D.; Robins, D.
#journal Nucleic Acids Res. (1986) 14:2539-2554
#title Sequence comparison of alleles of the fourth component of complement (C4) and sex-limited protein (Slp).
#cross-references MUID:86176748
#accession I48274
##status preliminary; translated from GB/EMBL/DBJ
##molecule_type mRNA
##residues 591-603, 'M', 605-1323, 'N', 1325-1452, 'V', 1454-1585, 'Q', 1587-1738 ##label RE2
##cross-references EMBL:X05314; NID:g50241; CDS_PID:g50242
REFERENCE
I54567
#authors Nonaka, M.; Nakayama, K.; Yeul, Y.D.; Shimizu, A.; Takahashi, M.
#journal Immunol. Rev. (1985) 87:81-99
#title Molecular cloning and characterization of complementary and genomic DNA clones for mouse C4 and Slp.
#cross-references MUID:86031969
#accession I54567
##status preliminary; translated from GB/EMBL/DBJ
##molecule_type DNA
##residues 1-128 ##label RE3
##cross-references GB:M12968; NID:g199267; CDS_PID:g199270
#accession I69023
##status preliminary; translated from GB/EMBL/DBJ
##residues 1724-1738 ##label RE4
##cross-references GB:M12969; NID:g199268; CDS_PID:g387439
COMMENT
This protein is synthesized as a single-chain precursor and, prior to secretion, is enzymatically cleaved to form a trimer of nonidentical chains (alpha, beta, and gamma) which are linked by disulfide bonds.
COMMENT
The activation of complement C4 by complement subcomponent C1s releases the C4a anaphylatoxin from the amino end of the alpha chain and generates C4b, which associates with the 2a fragment of complement factor 2 to form the classical-complement-pathway C3 convertase. The C4b, C2a fragment then associates with the 3b

fragment of complement factor 3 to form the classical-complement-pathway C5 convertase.

COMMENT C4a anaphylatoxin is a vasoactive peptide and a mediator of inflammation.

COMMENT The activity of C4b is regulated by proteolytic cleavage involving C4b-binding protein and factor I.

GENETICS

#introns 22/2; 86/3; 977/3; 1073/3; 1125/3; 1164/3; 1221/3

#note the list of introns is incomplete

CLASSIFICATION #superfamily alpha-2-macroglobulin

KEYWORDS acute phase; complement classical pathway; glycoprotein; hydrolase; inflammation; plasma; polymorphism; serine proteinase; thiolster bond

FEATURE

1-19 #domain signal sequence #status predicted #label SIG\

20-673 #product complement C4 beta chain #status predicted

#label BETA

240-673,754-1440,

1448-1738 #product complement C4b #status predicted #label C4B\

678-1440 #product complement C4 alpha chain #status predicted

#label ALP

678-753 #product C4a anaphylatoxin #status predicted #label C4A\

754-843 #region C4b-binding protein binding\

953-1332 #product C4d fragment #status predicted #label C4D\

1448-1738 #product complement C4 gamma chain #status predicted

#label GAM

224,743,1387 #binding site carbohydrate (Asn) (covalent) #status predicted\

753-754 #cleavage site Arg-Asn (complement subcomponent C1e) #status predicted\

1006-1009 #cross-link thiolester (Cys-Cln) #status predicted

SUMMARY #length 1738 #molecular-weight 192870 #checksum 4149

Query Match

Best Local Similarity 13.3%; Score 1605; DB 2; Length 1738;

Matches 449; Conservative 407; Mismatches 628; Indels 143; Gaps 111;

Db 137 ghivfvtqdpynpgqrvyrfaldqkmpstdfilitvsnshlrv-lkkel-ftats 194

Qy 124 GFLEIHTDKPVYTPDQSVKRVYSINDILKPAKRETVLTFIDPEGSEVDMEIDHIGII 183

Db 195 ifqdaftidpiseptgkwisarfedglesnrthfevkkyvlpnfekitpwkpyilmvp 254

Qy 184 SFPD-FKIPSNRYCMTIKNKYKEDFTTGTAFFENKVEYVLPHEVSIEP-E-YNFICY 240

Db 255 ensdeiqldiqarygkpv-gqvaytrfalnde--gqkrtflrletqaklvgrthie 311

Qy 241 KNFKNEFIIKARYNKVYKVTEDVYITFGIEDKDDQKEMQTAQNTALINGIAQVT 300

Db 312 iskdfqaaalkinigrvdeglrllyaataviespggemeeaeltsrffvssafsllder 371

Qy 301 FDSEATAVKELS--YYSELDANNKYLIYAVTVIESTGCFSEAEIPGKIKYPIKLNDA 358

Db 372 tkrhlvpqahfllqalvqmsgeasvnpvkva-tlvsgdsdqvlidqstnglqg-v- 428

Qy 359 TPLFLKPGIYP IKVQVQKSLDQVGVVPIVILNAQITDNNQETSD LDP SKSVTRVDDGVA 418

Db 429 sisfpipptvtelrlvsaagl-ypai--arltvqa-p-psrgtflaie-pldprsep 482

Qy 419 SEVLNIPSGVTVLEFNKTKDADLPENQARECYBATAYSSLSQSYLYIDWTDNHKALLV 478

Db 483 gdtfilnlgvqibaptfhyhmylierrgqimamg-rep-rtkv-tsavslvdhqlapsv 539

Qy 479 GEHNIIVTPKSPYDKITHYNYLILSKGKLIHFCTREKFSQASQINIPVTQNWPPSS 538

Db 540 yfayfvyhqq-h-p--vansalliniqsrdeqklqlkv-d-qakeyrnadmamlriqt-dsk 594

Qy 539 RLIVYIVTGEQTAEIUSDSVWLNIEEK-CGNQLQVHLSPDADAYSPGQTVSLNMTAGMD 597

Db 595 alvalgavdtalyavqgrehklpdmaskvfevinsyvcpgqgddalqvfdaglafad 654

Qy 598 SWVALAADVSAVYQORCAKPLE--RVFQFLEKSLDLCGAGGGLNANVFLAGLTLFT 655

Db 655 gdrltqtr-dlscpekkkrqkrnmfakvsekkgvsepdkrccqdgmtklpmkt 713

Qy 656 NANADDSOENDEPC-KEIL-RPRTL--QKKIEATAKYKHSVVKCCYDGAC-VNNDT 710

Db 714 ceqtaarvpqqa-crepfleccckfaedlrnqtrsqahlarhnmhmlqeedlideddliv 772

Qy 711 CQRAARISIGRCIKAFTECCVWASQIRANISHKDMQIGRL-H-M-K--TLIPVSKPEI 765

Db 773 rtsfpenwlrvepydskslltwpidsmttwelqhvslsksgklcvakptrvrvfrkh 832

Qy 766 RSTFESKMEVHLWPRKQLOFALPDSLITWEIQIGISNT-GICVADTVKAKVFRDVF 824

Db 833 lhlrlpisirrfefairpvlvnylnddravsvhvtpvgeclagggmmaqvtpvpsa 892

Qy 825 LEMNIPYSVVRGEQIQLKGTGVYNYRTSCMQFCVKMSAVEGICTSESPVHQGTSSKCV 884

Db 893 rp-vafsvptaaanv-plkvargv-fdl----g-davskilqiekagahreelv-yn 943

Qy 885 RQKVEGSSHLVTEVLPLEIGHNINFSLETWFGKEILVKTLRVVPEG-VKRESYSGVT 943

Db 944 ldp--l-nnlgitlpegsdpmivpdgdfeslvrtasepletmsegalspggvasll 1000

Qy 944 LDPGIVGTISRKEFFRIPDLDPVKTEIKRILSVKGLVGEIL-S-AVLSEGINILT 1001

Db 1001 rlpqqaetmiylaptltaenyldrteqsklsep-kd-havdliqk-gymziqqrk 1057

Qy 1002 HUPGSAEALMSVWPVYVHYLETGNHNIFHSDPLIEKQKLKKKKEGMSIMSRYN 1061

Db 1058 ndqsfqahrdstwtatfvklislacqevnspokloetaswllaq-qlqdgshdp 1116

Qy 1062 ADYSYVWKGSGASTMTAPALRVLGQVKNYQVQNSIGNSLMLVENYQLDNGSFKEN 1121

Db 1117 cpvlhramqggl-vgs-detvaltafvialhghldvfqddakqlknrveasitkansf 1174

Qy 1122 SQYQP IKLQGTLPVEARENSLYTAFVIGIRKA--F--DIC-PL-VKIDTALIKADNF 1174

Db 1175 lqkcasagllghaaitayaltlt-kasedlrnvahnsImamaeetqeh-lywglvlg 1232

Qy 1175 LLENTLPAQ-SFTTALSAVALSLGDKTHPQFASIV-SALKREALVKGNNPPIYR--F-W 1228

Db 1233 qdkvvlrptaprepvpqapalvietavall-hlliregkqkmadkaaswlthqgsf 1291

Qy 1229 KON--LQHKDSSVP-NTGT-AR-M-VETTAYALLSINIKD-I-NYVNPVIMLSEEQRY 1280

Db 1292 hqafstqdtvrtldaleaywiashtteekalkvrtlsemgrnglktghlhnqhvgkle 1351

Qy 1281 GGGFYSTQDTINAIEGLEYEY--SLLV--KQLSLMD-ID-VSYKHKA-LBNY--K-MT 1329

Db 1352 eelkfalgstievkvegnsgtklrlrtynvldmtnctcdqlgievkvtagvaywdane 1411

Qy 1330 DK-NFY-LGRPEVLL--ND-D-L-IVST-G-FG----S--GLAT-VHVTTVVHKT-STSE 1372

Db 1412 dyednydpmaaddpvpplqvtpqlqfegrrrrrreapkaeeesrvqvtvciwrnkd 1471

Qy 1373 EVCSEY-LKI-DTQDI--EA-SHYRGY-QN-SDYKR-I--VA----C-ASYKS-S-REESS 1416

Db	1707	qmcks	1711
Qy	1652	ttcss	1656

Search completed: Wed Jan 28 12:15:48 1998
Job time : 99 secs.

 WAPSPRL

 (TM)

Release 2.1D John F. Collins, Biocomputing Research Unit.
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MPsrch_pp protein - protein database search, using Smith-Waterman algorithm

Run on: Wed Jan 28 12:16:06 1998; MasPar time 37.99 Seconds
 935.704 Million cell updates/sec

Tabular output not generated.

Title: >US-08-487-283A-2
 Description: (1-1676) from US08487283A.pep
 Perfect Score: 12048
 Sequence: 1 MGLLGILCLFLFGKTMQGE.....CQAFIANLDEPAEDIFLNGC 1676

Scoring table: PAM 150

Gap 11

Searched: 59021 seqs, 21210388 residues

Post-processing: Minimum Match 0%
 Listing first 45 summaries

Database: swiss-prot34
 1:part1 2:part2 3:part3 4:part4 5:part5 6:part6 7:part7
 8:part8 9:part9 10:part10 11:part11

Statistics: Mean 58.177; Variance 98.799; scale 0.589

Pred. No. is the number of results predicted by chance to have a
 score greater than or equal to the score of the result being printed,
 and is derived by analysis of the total score distribution.

SUMMARIES

Result	No.	Score	Match	Length	ID	Description	Pred. No.
1	12048	100.0	1676	2	CO5_HUMAN	COMPLEMENT C5 PRECURS	0.00e+00
2	9789	81.3	1680	2	CO5_MOUSE	COMPLEMENT C5 PRECURS	0.00e+00
3	2367	19.6	1651	2	CO3_NAJANA	COMPLEMENT C3 PRECURS	0.00e+00
4	2361	19.6	1663	2	CO3_HUMAN	COMPLEMENT C3 PRECURS	0.00e+00
5	2309	19.2	1673	2	CO3_LAMJA	COMPLEMENT C3 PRECURS	0.00e+00
6	2296	19.1	1663	2	CO3_MOUSE	COMPLEMENT C3 PRECURS	0.00e+00
7	2302	19.1	1666	2	CO3_CAVPO	COMPLEMENT C3 PRECURS	0.00e+00
8	2255	18.7	1663	2	CO3_RAT	COMPLEMENT C3 PRECURS	0.00e+00
9	2106	17.5	1620	2	CO3_EPTBU	COMPLEMENT C3 (CONTAI	0.00e+00
10	2002	16.6	1640	2	CO3_ONCHY	COMPLEMENT C3-1 (CONT	0.00e+00
11	1723	14.3	1741	2	CO4_HUMAN	COMPLEMENT C4 PRECURS	0.00e+00
12	1605	13.3	1738	2	CO4_MOUSE	COMPLEMENT C4 PRECURS	0.00e+00
13	796	6.6	726	2	CO3_RABIT	COMPLEMENT C3 ALPHA C	5.45e-148

14	670	5.6	1477	1	AI13_RAT	ALPHA-1-INHIBITOR III	2.99e-118
15	661	5.5	1476	1	A2MG_MOUSE	MURINOGLOBULIN 1 PREC	3.83e-116
16	552	4.8	1474	1	A2MG_HUMAN	ALPHA-2-MACROGLOBULIN	8.97e-98
17	581	4.8	1482	8	P2P_HUMAN	PREGNANCY ZONE PROTEI	1.53e-97
18	508	4.2	323	2	CO3_XENIA	COMPLEMENT C3 (FRAGME	8.31e-81
19	507	4.2	1472	1	A2MG_RAT	ALPHA-2-MACROGLOBULIN	1.40e-80
20	502	4.2	1473	7	OVO5_CHICK	OVOSTATIN PRECURSOR (1.92e-79
21	493	4.1	1451	1	A2MH_MOUSE	MURINOGLOBULIN 2 PREC	2.11e-77
22	399	3.3	74	2	CO5A_PIG	COMPLEMENT C5A ANAPHY	2.09e-56
23	385	3.2	74	2	CO5A_BOVIN	COMPLEMENT C5A ANAPHY	2.44e-53
24	351	2.9	76	2	CO5A_RAT	COMPLEMENT C5A ANAPHY	5.68e-46
25	171	1.4	76	2	CO4A_RAT	COMPLEMENT C4A ANAPHY	3.12e-10
26	170	1.4	77	2	CO4A_BOVIN	COMPLEMENT C4A ANAPHY	4.64e-10
27	139	1.2	77	2	CO3A_PIG	COMPLEMENT C3A ANAPHY	5.44e-05
28	121	1.0	798	10	YA73_SCHPO	HYPOTHETICAL 92.1 KD	2.39e-02
29	115	1.0	1292	8	RPOC_MYCGE	DNA-DIRECTED RNA POLY	1.57e-01
30	105	0.9	257	11	YGBI_HAEIN	HYPOTHETICAL TRANSCRI	3.00e+00
31	106	0.9	270	11	YD1J_SCHPO	HYPOTHETICAL 31.5 KD	2.26e+00
32	105	0.9	328	1	ASCD_YERPS	CDP-6-DEOXY-DELTA-3,4	3.00e+00
33	110	0.9	341	11	YEJK_HAEIN	HYPOTHETICAL PROTEIN	7.10e-01
34	108	0.9	461	1	AD4B_BOVIN	STEROID HORMONE RECEP	1.27e+00
35	106	0.9	462	1	AD4B_MOUSE	STEROID HORMONE RECEP	2.26e+00
36	112	0.9	472	8	PSBC_SINY3	PHOTOSYSTEM II 44 KD	3.91e-01
37	108	0.9	589	5	KPYK_LACDE	PYRUVATE KINASE (EC 2	1.27e+00
38	108	0.9	612	10	UNC6_CAEEL	UNC-6 PROTEIN PRECURS	1.27e+00
39	108	0.9	943	5	IROA_NEIME	IRON-REGULATED OUTER	1.27e+00
40	113	0.9	1024	10	UBA1_YEAST	UBIQUITIN-ACTIVATING	2.90e-01
41	106	0.9	1030	10	VPPI_CAEEL	PUTATIVE CLATHRIN-COA	2.26e+00
42	112	0.9	1517	4	GLTB_ECOLI	GLUTAMATE SYNTHASE (N	3.91e-01
43	107	0.9	1736	11	ZO1_HUMAN	TIGHT JUNCTION PROTEIN	1.70e+00
44	108	0.9	1805	10	Y218_MYCGE	HYPOTHETICAL PROTEIN	1.27e+00
45	106	0.9	5147	3	FAT_DROME	CADHERIN-RELATED TUMO	2.26e+00

ALIGNMENTS

RESULT	1	CO5_HUMAN	STANDARD;	PRT;	1676 AA.
ID	CO5_HUMAN				
AC	P01031;				
DT	21-JUL-1986	(REL. 01, CREATED)			
DT	01-DEC-1992	(REL. 24, LAST SEQUENCE UPDATE)			
DT	01-FEB-1996	(REL. 33, LAST ANNOTATION UPDATE)			
DE	COMPLEMENT C5 PRECURSOR (CONTAINS: C5A ANAPHYLATOXIN).				
GN	C5.				
OS	HOMO SAPIENS (HUMAN).				
OC	EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA;				
OC	EUTHERIA; PRIMATES.				
RN	[1]				
RP	SEQUENCE FROM N.A.				
RX	MEDLINE; 91079575.				
RA	HAVILAND D.L., HAVILAND J.C., FLEISCHER D.T., HUNT A., WETSEL R.A.;				
RL	J. IMMUNOL. 146:362-368 (1991).				
RN	[2]				
RP	SEQUENCE OF 412-1676 FROM N.A.				
RX	MEDLINE; 88209511.				
RA	WETSEL R.A., LEMONS R.S., LEBEAU M.M., BARNUM S.R., NOACK D.,				
RA	TACK B.F.;				
RL	BIOCHEMISTRY 27:1474-1482 (1988).				
RN	[3]				
RP	SEQUENCE OF 412-902 FROM N.A.				
RX	MEDLINE; 85130937.				
RA	LUNDWALL A.B., WETSEL R.A., KRISTENSEN T., WHITEHEAD A.S.,				
RA	WOODS D.E., OGDEN R.C., COLTEN H.R., TACK B.F.;				
RL	J. BIOL. CHEM. 260:2108-2112 (1985).				

RP SEQUENCE OF 678-751.
RX MEDLINE; 79005687.
RA FERNANDEZ H.N., HUGLI T.E.;
RL J. BIOL. CHEM. 253:6955-6964 (1978).
[5]
RP SEQUENCE OF 678-751 FROM N.A.
RX MEDLINE; 91144547.
RA BOHNSACK J.F., MOLLISON K.W., BUKO A.M., ASHWORTH J.C., HILL H.R.;
RL BIOCHEM. J. 273:635-640 (1991).
[6]
RP STRUCTURE BY NMR OF CSA.
RX MEDLINE; 88309754.
RA ZUIDERWEG E.R., MOLLISON K.W., HEKIN J., CARTER G.W.;
RL BIOCHEMISTRY 27:3568-3580 (1988).
[7]
RP STRUCTURE BY NMR OF CSA.
RX MEDLINE; 89207527.
RA ZUIDERWEG E.R., NETTESHEIM D.G., MOLLISON K.W., CARTER G.W.;
RL BIOCHEMISTRY 28:172-185 (1989).
[8]
RP STRUCTURE BY NMR OF CSA.
RX MEDLINE; 89274164.
RA ZUIDERWEG E.R., FESIK S.W.;
RL BIOCHEMISTRY 28:2387-2391 (1989).
CC -1- FUNCTION: ACTIVATION OF C5 BY A C5 CONVERTASE INITIATES THE
CC SPONTANEOUS ASSEMBLY OF THE LATE COMPLEMENT COMPONENTS, C5-C9,
CC INTO THE MEMBRANE ATTACK COMPLEX. C5B HAS A TRANSIENT BINDING SITE
CC FOR C6. THE C5B-C6 COMPLEX IS THE FOUNDATION UPON WHICH THE LYtic
CC SUBUNIT IS ASSEMBLED.
CC -1- SUBUNIT: C5 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 BASIC
CC RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY A DISULFIDE
CC BOND. C5 CONVERTASE ACTIVATES C5 BY CLEAVING THE ALPHA CHAIN,
CC RELEASING C5A ANAPHYLATOXIN & GENERATING C5B (BETA CHAIN + ALPHA
CC CHAIN).
CC -1- SIMILARITY: TO C3, C4 AND ALPHA-2-MACROGLOBULIN.
CC -1- FUNCTION: DERIVED FROM PROTEOLYTIC DEGRADATION OF COMPLEMENT C5,
CC C5 ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS. IT
CC INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR
CC PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND
CC BASOPHILIC LEUKOCYTES. C5A ALSO STIMULATES THE LOCOMOTION OF
CC POLYMORPHONUCLEAR LEUKOCYTES (CHEMOKINESIS) AND DIRECT THEIR
CC MIGRATION TOWARD SITES OF INFLAMMATION (CHEMOTAXIS).
CC -1- CAUTION: REF. 3 SEQUENCE DIFFERS FROM THAT SHOWN FROM POSITION 855
CC ONWARD DUE TO THE PRESENCE OF AN ALU REPEAT.
CC -1- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN.
DR EMBL; M57729; G179983; --.
DR EMBL; M65134; G179692; --.
DR PIR; S15121; S15121...
DR HSSP; P01032; 1C5A.
DR MIM; 120900; --.
KW PROSITE; PS00477; ALPHA 2 MACROGLOBULIN.
KW COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; GLYCOPROTEIN;
KW PLASMA; MEMBRANE ATTACK COMPLEX; CYTOLYSIS; INFLAMMATORY RESPONSE;
KW SIGNAL; POLYMORPHISM.
FT SIGNAL 1 18 POTENTIAL.
FT CHAIN 19 673 COMPLEMENT C5 BETA CHAIN.
FT PROPEP 674 677
FT CHAIN 678 1676 COMPLEMENT C5 ALPHA CHAIN.
FT PEPTIDE 678 751 C5A ANAPHYLATOXIN.
FT CHAIN 752 1676 C5B (ALPHA').
FT DOMAIN 698 732 ANAPHYLATOXIN-LIKE.
FT DISULFID 698 724

FT DISULFID 699 731
FT DISULFID 711 732
FT CARBOHYD 741 741
FT CARBOHYD 911 911 POTENTIAL.
FT CARBOHYD 1115 1115 POTENTIAL.
FT CARBOHYD 1630 1630 POTENTIAL.
FT VARIANT 518 518 F -> S.
SQ SEQUENCE 1676 AA; 188331 MW; 9D5C6E59 CRC32;
Query Match 100.0%; Score 12048; DB 2; Length 1676;
Best Local Similarity 100.0%; Pred. No. 0.00e+00;
Matches 1676; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Db 1 mglilicfliflgtwgeqyvisapkiirvgaseniivqygyteafdatiskayp 60
Qy 1 MGLLIGLILFLIFLGKWTGQEQTYVISAPKIFRVRGASENIIVQYGYTEAFDATISIKSYP 60
Db 61 dkfkfssghvhlssenkfmsaalltiqpkqpqqgnpvyvylevskhfkskmpit 120
Qy 61 DKFKFSYSSGHVHLSSENKFNQNGAILTIQPKQLPGQNPVSYVYLEVSKHFKSKRMPIIT 120
Db 121 ydnofifihitdkpytpdgskvrvyslnddlkpkakrtetvltfdpegsevmveidhi 180
Qy 121 YDNQFLFIHTDKPYTPDQSVKVRVYSLNDDLPKAKRTETVLTFDPEGSEVMVEIDHI 180
Db 181 giisfpdkipenrygmwtikakykedfstgtayfevkeyvlpfhfsieieynfigy 240
Qy 181 GIISFPDKIPSNPRYGMWTIKAKYKEDFSTGTAYFEVKEYVLPFHFSVSIEPEYNFIGY 240
Db 241 knfkneitkaryfynkvvteadvitfgiredlkddqkmmqtamqntmlinglaqt 300
Qy 241 KNFKNEITIKARYFNKVVTADVITFGIREDLKDDQKEMMQTAMQNTMLINGLAQVT 300
Db 301 fsetavkelsyysledlnmkyllylvtviestgqfseaeipgikyvlsykylnlvatp 360
Qy 301 FSETAVKELSYYSLEDLNKYLVIYVTVIESTGQFSEAEIPGIRYVLSPIYKINLVATP 360
Db 361 lflkpgipypikvqkdsldqlvggvpiinaactidvncqetadldpsksvtrvddgvasf 420
Qy 361 LFLKPGIPYPKVVQKDSLDQLVGGVPILNAQTTIDVNCQETSDLPDSKSVTRVDDGVASF 420
Db 421 vlnlpesgvtvlefntvdapdlpeengaregyraiysslesqylyidwtdhkaallvge 480
Qy 421 VLNLPESGVTVLEFNVTDAVDLPENQAREGYRAIYSSLSQSYLYIDWTDNHKALLVGE 480
Db 481 hlniivtpkspydikithynylilekgkiihfgtrekfsdaayqsiinipvtqmvpsrl 540
Qy 481 HLNIIIVTPKSPYDIKITHYNYLILSKGKIIHFGTREKFSDSYQSINIPVTQNMVPSRSL 540
Db 541 lvyvitytgeqtaelvsdsvlnieekcngqlqvhlpdadavspqgtvsiinmatgmsdw 600
Qy 541 LVYVITYTGEQTAELVSDSVLNIEEKCGNQLQVHLPDADAVSPQGTVSIINMATGMSDW 600
Db 601 alaadaevygvqrakklervfglekediqcgaggglnnanvfhlagltfltnanad 660
Qy 601 ALAADADEVYGVQRKKLERVFGLEKEDIQCGAGGGLNNANVFHLAGLTFLTNANAD 660
Db 661 daqendepckelprtrtlqkkieiaakysvkkccydcacvmdetceqraarisl 720
Qy 661 DSQENDEPCKELPRTRTLQKKEIEIAAKYKHSVVKCCYDCACVNMDETCEQRAARISL 720
Db 721 gprcikafteccvvaaglraniashkdmqlgrlhmktllpvskeirsyfswlwehvlv 780
Qy 721 GPRCIKAFTECCVVASQLRANISHKDMQLGRLHMKTLLPVSKEIRSYFFESWLWEHVLV 780

Db 781 prtkqlfaldtsitweiqigisntgicvadtkvfkdvflemnipyvvrgeqiq 840
|||||
Qy 781 PRKQQLFALDTSITWEIQIGISNTGICVADTKVFKDVFLEMNIPYVVRGEQIQ 840
Db 841 lkgvnyvrtsgmfcvktasavegictseepvldhgtkaskcvrqkvegssahlvttv 900
|||||
Qy 841 LKGTVNYRTSGMFCVKTASAVEGICTSESPVIDHQTKSKCRQKVEGSSSHLVTTV 900
Db 901 lpleighninfletwfkellvktlrvvpegvkresygvrtldprgiygtisrrkefp 960
|||||
Qy 901 LPLEIGHNINFLETWFKELLVKTLRVVPEGVKRESYGVRTLDPRGITYGTISRRKEFP 960
Db 961 yripldvpkteikrslvkvglvgeilaevisqeginlthlpkgaeealmsvvpvyf 1020
|||||
Qy 961 YRIPLDVPKTEIKRSLVKVGLVGEILAEVISQEGINLTHLPKGAEEALMSVVPVYF 1020
Db 1021 vfhyletgnhwnifhscdpliekqlkkklkegmlelmsvrynadysvkwkggsastwita 1080
|||||
Qy 1021 VFHYLETGNHWNIFHSDPLIEKQKAKKKLKEGMLELMSVRNADYSVWMKGGASWTITA 1080
Db 1081 falrvlgvmkyvegmqmsicnllwvnyqldngsfkenskqppiklqgtlpvearen 1140
|||||
Qy 1081 FALRVLGVMKYVEGMQMSICNLSLWVNYQLDNGSFKENSKQPPIKLQGTLPVEAREN 1140
Db 1141 slyltaftvigrkafdcipvlkicplvkdialkadhflentlpaqstflaisayalslqdk 1200
|||||
Qy 1141 SLYLTAFTVIGRKAFDCIPVLKICPLVKDIALKADHFLENTLPAQSTFLAISAYALSQDK 1200
Db 1201 thpofrsivalkrealvkgppiyrfwkhldkhdssvntgtarmvettayalltsln 1260
|||||
Qy 1201 THPOFRSIVALKREALVKGPPIYRFWKHLDKHDSSVNTGTARMVETAYALLTSIN 1260
Db 1261 lkdinvmpkvlseeqyvggfygtdtinaeglteysllvkqlrlmdldvsvykhk 1320
|||||
Qy 1261 LKDINVMPKVLSEEQYVGGFYSTQDTINAEGLTEYSLLVKQLRLMDLDVSVYKHK 1320
Db 1321 galhnykmtkdnflgrpvevllnddlivstfgsglatvhtvtvvhktstseevcsfvlk 1380
|||||
Qy 1321 GALHNYKMTKDNFLGRPVEVLLNDLIVSTFGSGLATVHTVTVVHKTSTSEEVCSFVLK 1380
Db 1381 idtdieahyrgyngsdykrivacasykpsreessgshavmdielptgisaneedlk 1440
|||||
Qy 1381 IDTDIEAHYRGYNGSDYKRIVACASYKPSREESSGSHAVMDISLPTGISANEEDLK 1440
Db 1441 alvegvdqlfcdyqikdghvilqinspsdflcvrfrifelfevgflspatftvyeyhr 1500
|||||
Qy 1441 ALVEGVVDQLFCDYQIKDGHVILQINSPSDFLCVRFRIFELFEVGFLESPATFTVYEYHR 1500
Db 1501 pdkqctmfystemkikqkveagackveadgmgqeeldltiaeatriktackpelaya 1560
|||||
Qy 1501 PDKQCTMFYSTEMKIKQKVEAGACKVEADGGMQEELDLTIAEATRIKTKACKPELAYA 1560
Db 1561 ykveiteltvenfvkykatlldiyltgeavaekdseiftikvctcnaelvkgrqylm 1620
|||||
Qy 1561 YKVEITELTVENFVKYKATLLDIYLTGEAVALKDESEIFTIKVCTCNAELVKGRQYLM 1620
Db 1621 gkealqikynfsfrylpldsaltwiewprdtccsqcqlanldefaediifngc 1676
|||||
Qy 1621 GKEALQIKYNFSFRYLPPLDSALTWIEWPRDTCCSQCAFANLDEFAEDIIFNGC 1676

RESULT 2

ID CO5_MOUSE STANDARD; PRT; 1680 AA.

AC P06684;
DT 01-JAN-1988 (REL. 06, CREATED)
DT 01-NOV-1990 (REL. 16, LAST SEQUENCE UPDATE)
DT 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE)
DE COMPLEMENT C5 PRECURSOR (CONTAINS: C5A ANAPHYLATOXIN).
GN C5.
OS MUS MUSCULUS (MOUSE).
OC EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA;
OC EUThERIA; RODENTIA.
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE; 90153853.
RA WETSEL R.A., FLEISCHER D.T., HAVILAND D.L.;
RL J. BIOL. CHEM. 265:2435-2440 (1990).
RN [2]
RP SEQUENCE OF 41-1680 FROM N.A.
RX MEDLINE; 87185363.
RA WETSEL R.A., OGATA R.T., TACK B.F.;
RL BIOCHEMISTRY 26:737-743 (1987).
CC -!- FUNCTION: ACTIVATION OF C5 BY A C5 CONVERTASE INITIATES THE
CC SPONTANEOUS ASSEMBLY OF THE LATE COMPLEMENT COMPONENTS, C5-C9,
CC INTO THE MEMBRANE ATTACK COMPLEX. C5B HAS A TRANSIENT BINDING SITE
CC FOR C6. THE C5B-C6 COMPLEX IS THE FOUNDATION UPON WHICH THE LYtic
CC COMPLEX IS ASSEMBLED.
CC -!- SUBUNIT: C5 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 BASIC
CC RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY A DISULFIDE
CC BOND. C5 CONVERTASE ACTIVATES C5 BY CLEAVING THE ALPHA CHAIN,
CC RELEASING C5A ANAPHYLATOXIN & GENERATING C5B (BETA CHAIN + ALPHA'
CC CHAIN).
CC -!- SIMILARITY: TO C3, C4 AND ALPHA-2-MACROGLOBULIN.
CC -!- FUNCTION: DERIVED FROM PROTEOLYTIC DEGRADATION OF COMPLEMENT C5,
CC C5 ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS. IT
CC INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR
CC PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND
CC BASOPHILIC LEUKOCYTES. C5A ALSO STIMULATES THE LOCOMOTION OF
CC POLYMORPHONUCLEAR LEUKOCYTES (CHEMOKINESIS) AND DIRECT THEIR
CC MIGRATION TOWARD SITES OF INFLAMMATION (CHEMOTAXIS).
CC -!- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN.
EMBL; M35525; G309124; -.
DR EMBL; M35526; G309123; -.
DR PIR; A27538; A27538.
DR PIR; A35530; A35530.
DR HSP; P01032; 1C5A.
DR PROSITE; PS00477; ALPHA 2 MACROGLOBULIN.
KW COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; GLYCOPROTEIN;
KW PLASMA; MEMBRANE ATTACK COMPLEX; CYTOLYSIS; INFLAMMATORY RESPONSE;
KW SIGNAL.
FT SIGNAL 1 18
FT CHAIN 19 1680 COMPLEMENT C5.
FT CHAIN 19 674 COMPLEMENT C5 BETA CHAIN.
FT PROPEP 675 678
FT CHAIN 679 1680 COMPLEMENT C5 ALPHA CHAIN.
FT PEPTIDE 679 755 C5A ANAPHYLATOXIN.
FT CHAIN 756 1680 C5B (ALPHA').
FT DOMAIN 702 736 ANAPHYLATOXIN-LIKE.
FT DISULFID 702 728 BY SIMILARITY.
FT DISULFID 703 735 BY SIMILARITY.
FT DISULFID 715 736 BY SIMILARITY.
FT CARBOHYD 427 427 POTENTIAL.
FT CARBOHYD 915 915 POTENTIAL.
FT CARBOHYD 1119 1119 POTENTIAL.
FT CARBOHYD 1633 1633 POTENTIAL.
FT VARIANT 216 216 Y -> L (IN DEFECTIVE VARIANT C5D).
FT VARIANT 217 1680 MISSING (IN DEFECTIVE VARIANT C5D).

SQL	SEQUENCE	1680 AA; 18887 MM; AA170448 CRC32;
	Query Match	81.3%; Score 9789; DB 2; Length 1680;
	Best Local Similarity	77.8%; Pred. No. 0.00e+00;
	Matches 1307; Conservative	218; Mismatches 149; Indels 7; Gaps
Db	1	mglwgllcllflldtwwqetvysaapkllrvsgsenvviqhyqteafdatlkeyp 60
Qy	1	MGLLGLLCLFLFLGKTKQETVVSAPKIFRVGASENIVIQVGYTEAFDATISIKSY 60
Db	61	dkkvtfssgynspenkfnaalltqlnqvprespsvshvylevskhfekskpit 120
Qy	61	DKKTSYSGGHVLSSENKFNQNSALLTQPKQLGCGQNPVSYVYLEVSKHFESKRW 120
Db	121	ynngilfhcdkpytpdqgcvkiryvsigddlkapkretvltfidpegsevd 180
Qy	121	YNGGLFIHTDKPVTDPQSVKRVVYSLNDJAPAKRETVLTFIDPEGSEVDMVEE 180
Db	181	qiiefpdkipsnpgywtikanykdkftttatgafelkeylprfsvsie 240
Qy	181	GIISFPDKIPSNRPGWMTIKAKYKEDFTTGTAFFEYKEYLPHFSVSIENFY 240
Db	241	kfnkfnfeitkaryfynkvpdaevyafgliredildekqmmhkatqaaklvdv 300
Qy	241	KFNKFNFEITIKARYFYNKVTEADVYITFGIREDLKDDQEMMQATQNTMLING 300
Db	301	fdsetavkelsyns ledlnmkylyiavtveseggfseeaipygvkylspt 360
Qy	301	FDESETAVKELSYSLDNLNNKYLYIAVTVIESTGFSSEAEIPGIVKYLSPYK 360
Db	361	lvfkpgipfsikaqvkdelegavgyvpytlnaqtvdvmetdsletkrsith 420
Qy	361	LFVFKPGIPYPKQVQKSLDQLVGGVPVLNQAQTIQVNETSDLDPSKSVTV 420
Db	421	vlnlpsnvtlwfektldpelpenqaskeyeavyslsqsviyiawteny 480
Qy	421	VNLNPSGVTVLEFENVKTDADLPENQAREGYRAIAYSLSQSYLYIDWTDH 480
Db	481	ylnlmvtptsepvydkithynylilekqivqygtrekfssatymini 540
Qy	481	HLNIVTPKSPYIDKITHYNLYLSKGIHFGTRESKSDASYSINIPVTQNM 540
Db	541	lvyyivrtgectaelvadavwinieekcnqlqvlhlpdeyvyvyspg 600
Qy	541	LVYYIVTGTQAEIYSDSVWLNTEEEKCNQLQVHLSPDADAYSQGTVSLN 600
Db	601	aleadvravykvgnakramrvqfaldekedlgcagagghdnadvflag 660
Qy	601	ALAAVDSAVYGVQRCAPKPLERVFQFL-EKSDLCCGAGGLNANNVFLAG 659
Db	661	ddshvrdscelskzrnllhllrkieegaakykhsvpkcccydgarnvf 720
Qy	660	DDSGNDEPCKEILRPRTQL--K-KIEIAKYKUSVWVKCCYDGCANNDE 716
Db	721	rvtigpiclrafneccitankirspkpvqlqrihiktllpvmkadirs 780
Qy	717	RISLGPICIAFTECCVWASQLRANISHKQMLGRUHKMTLLPYSKPEIR 776
Db	781	lhrvpkrkqlqvtlpdalttweiqqigisdnqicvadtlkakvfkve 840
Qy	777	VHLVPRKQLQALPDSLTTWEIQIGISNTGICVADTVKAKVFKOVLENN 836
Db	841	eqiqlkgtyvnymtsgtkfcvkmsavegitsgssaaalhtcrsprscv 900

Qy	837	EQIKGKTGVTNVRTSGMQFCVKMSAVEGICTSESPVIDHOGTKSKCVRQKVEGSSSHLV	896
Db	901	tftllpleighheinfelsfgkdilvklrlrwpvgkresyagvilpdkpgirgvmnr	960
Qy	897	TFVLPLEIGLJHNFISLETWFGEILVKTLRVPEGVKRESYSGVTLDPRGITGTSIRR	956
Db	961	kafpyripldvpkktkverilevkgllvgfeflvtakeginilthlpkgsaaelmaia	1020
Qy	957	KEFPYRIPDLVPKTEIKRILSVKGLLVGEILSAVLSQEGINILTHLPKGSAAELMSV	1016
Db	1021	pvfyvfyhleaqnwnfiydpdtlskrgelekkikgqvsvmsyrnadyswmkqasast	1080
Qy	1017	PVFYVYHLETGNHNHNFHSDDLIEKQKIKKKLKEGMLSTMSYRNADYSYVWKGCAS	1076
Db	1081	wltafalrvlqqvakvqkqdensicnsllwvekcqlengsfkensksgylpklqgtlpae	1140
Qy	1077	WUTAFALRVLQGVKNYVEQNGQNSICNSLWLWLVENTQJLNGSFRKNSQJPIKUQGTPE	1136
Db	1141	aqetllvtafavigirkvadicmtkhtaldkadefflentlpstetftlaiavays	1200
Qy	1137	ARENSLTLAFTVIGIRKAPDLCPLVIRKIDTALIKADNFLEENTLPAOSTFTLAIAYALS	1196
Db	1201	lqdrthprfrlivaalkaefvkgdppiyryrdtklrpdsvpasgtagmvettavall	1260
Qy	1197	LGDKTHPQFRISVALKREALVKGNPPYIRFKNDLQIHKDSSVPNTGTARWVETAYALL	1256
Db	1261	aaclkdmnyampiikweeqryggfyetqdtinaieqlteyallklqihldmndinva	1320
Qy	1257	TSJNLKIDINYNVPKIVLSEQRYSGGFYSTQDTINALEGLTEYSLVWKQLRLSMDIDVS	1316
Db	1321	ykhqgdhkykvtexhflgrpvevslnddlrvstgysglatvyvktvwhkjsvseeecs	1380
Qy	1317	YKHGALHNYKMTDKNPLGRPVEVLNDDLIVSTGCGSLATVYVTVVWTKTSTEEVCS	1376
Db	1381	fylkidtdieashfr-lsdsgfkriiacasykpskeestsgshavmdislptqigan	1439
Qy	1377	FYIKIDTQDTEAS-HYRGVNSDVKRIVACASYKPSREESSGSSHAVMDISLPTGISAN	1435
Db	1440	eedlralvevgdqlttdyqikghvilqinslpsrddflcvrfeifqvgfnpattftv	1499
Qy	1436	EEDLALVEGVQDLFTDYQIKOGHVILQINSIPSSDFLCVRFRIFELFEVGLSPATFTV	1495
Db	1500	veyhrpdqctmisyiedtrlkqvcgaactcveadcaqlqaevdlaisadrskeackp	1559
Qy	1496	YEYHRPDQCTMFYSTNIIQKVCEGAACKVEADCGQMEELDOLTISAETRKQTACKP	1555
Db	1560	etayaykvritateenfvkyatatllyvtktgea--adensevtfikmactnanlvkgk	1618
Qy	1556	EIAYAYKVSTSTIVENVFKYKATLIDYKTCGEAVAEKDSIEFTIKKV7CINAEUVGR	1615
Db	1619	qylimgkevlqikhnfkyipyidsatwiewpdttdtccpsccaqfvenlmmfaedflns	1678
Qy	1616	QYILMGKEALQIKNFSTRYYIPJDSLTWIEWPDRDTCCSCQAFLANLDEFAEDFLNG	1675
Db	1679	c	1679
Qy	1676	c	1676
RESULT 3			
ID	CO3.NAJNA	STANDARD;	PRT; 1651 AA.
AC	Q01833;		
DT	01-JUL-1993	(REL. 26, CREATED)	
UDT	01-JUL-1993	(REL. 26, LAST SEQUENCE UPDATE)	

DT 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE)
DE COMPLEMENT C3 PRECURSOR (CONTAINS: C3A ANAPHYLATOXIN).
GN C3.
OS NAJA NAJA (INDIAN COBRA).
OC EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; REPTILIA;
OC LEPIDOSAURIA; SERPENTES.
RN [1]
RV SEQUENCE FROM N.A.
RP TISSUE=LIVER;
RC MEDLINE; 93056528.
RA FRITZINGER D.C., CONNELLY M., PETRELIA E.C., BREDEHORST R.,
RL J. IMMUNOL. 149:3554-3562(1992).
CC -!- FUNCTION: C3 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE
CC COMPLEMENT SYSTEM. ITS PROCESSING BY C3 CONVERTASE IS THE CENTRAL
CC REACTION IN BOTH CLASSICAL AND ALTERNATIVE COMPLEMENT PATHWAYS.
CC AFTER ACTIVATION C3B CAN BIND COVALENTLY, VIA ITS REACTIVE
CC THIOLESTER, TO CELL SURFACE CARBOHYDRATES OR IMMUNE AGGREGATES.
CC -!- SUBUNIT: C3 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 ARG
CC RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY A DISULFIDE
CC BOND. C3 CONVERTASE ACTIVATES C3 BY CLEAVING THE ALPHA CHAIN,
CC RELEASING C3A ANAPHYLATOXIN & GENERATING C3B (BETA CHAIN + ALPHA
CC CHAIN).
CC -!- SIMILARITY: TO C4, C5 AND ALPHA-2-MACROGLOBULIN.
CC -!- FUNCTION: DERIVED FROM PROTEOLYTIC DEGRADATION OF COMPLEMENT C3.
CC C3A ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS. IT
CC INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR
CC PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND
CC BASOPHILIC LEUKOCYTES.
CC -!- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN.
DR EMBL; L02365; G213373; --.
DR PIR; A46513; A46513.
DR HSP; P01032; IC5A.
DR PROSITE; P500477; ALPHA 2 MACROGLOBULIN.
KW COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; PLASMA;
KW INFLAMMATORY RESPONSE; SIGNAL.
FT SIGNAL 1 22
FT CHAIN 23 1651 COMPLEMENT C3.
FT CHAIN 23 655 BETA CHAIN.
FT CHAIN 661 1651 ALPHA CHAIN.
FT PEPTIDE 661 738 C3A ANAPHYLATOXIN.
FT CHAIN 739 1651 C3B (ALPHA' CHAIN).
FT SITE 738 739 CLEAVAGE (BY C3 CONVERTASE).
FT DOMAIN 683 718 ANAPHYLATOXIN-LIKE.
FT DISULFID 546 807 INTERCHAIN (BY SIMILARITY).
FT DISULFID 615 650 BY SIMILARITY.
FT DISULFID 683 710 BY SIMILARITY.
FT DISULFID 684 717 BY SIMILARITY.
FT DISULFID 697 718 BY SIMILARITY.
FT DISULFID 863 1501 BY SIMILARITY.
FT DISULFID 1091 1147 BY SIMILARITY.
FT DISULFID 1346 1477 BY SIMILARITY.
FT DISULFID 1377 1446 BY SIMILARITY.
FT DISULFID 1494 1499 BY SIMILARITY.
FT DISULFID 1506 1578 BY SIMILARITY.
FT DISULFID 1525 1649 BY SIMILARITY.
FT THIOLEST 999 1002 BY SIMILARITY.
SQ SEQUENCE 1651 AA; 184926 MW; 2958575F CRC32;

Query Match 19.6%; Score 2367; DB 2; Length 1651;
Best Local Similarity. 30.2%; Pred. No. 0.00c+00;
Matches 518; Conservative 416; Mismatches 655; Indels 127; Gaps 97;

DB 4 malyvlaalligfgg-s-shgalytlitpavlrtdteeqilveahgdgtpkldifvhd 61

Qy 1 MGLLGILCFLLI-FLGKTMGQEQTYVISAPKIFRVGASENIVQVYCYTEAFDATISIKSY 59
Db 62 prkqhtlfgervdmqagmfvtpti-kvpakehndsknqvvvkvtpgqvalekvvl 120
Qy 60 PDKKFSYSSGHVLS-SENKFNQSAIITIQKQLPGQNPVSVYVLEVSKHFSKSRMP 118
Db 121 leyqsgfviqtdkgytpqspvrvsvfvdhnmhmdktviveftqpegivsvs-ekpm 179
Qy 119 ITYONGLELIHTDKPVYTPDQSVKRVYSINDDIKPAKRETVLTFIDPEGSEVDWVEID 178
Db 180 psgsir-p-ynlpaelvsfqtwkavakyspeesyayfvdyreyvlpsfvevrlpsdkfl 237
Qy 179 HIGIISPFDFKIPSNPRYGMWTKAKYKREDEFTTGTAYFEVKYVLPHFVSIEPEYNF 238
Db 238 -yidnknfhveitarylgykkv-egvafvfgvk-i-ddakkeipsdeltipiidag 292
Qy 239 GYKN-FNFEITIKARYFNKRVVTEADVYITFGIREDLKDDQKEMQMTAMQNTMLINGIA 297
Db 293 eatikrdt-lrs-rfdlnqlvghtlysvvtitesgdmvvtteggihivtpeyqiyft 350
Qy 298 QVTFDSETAVKELSYSLIEDLANKVLYIAVTVIESTGFSSEAEIPGKYVLSPYKLV 357
Db 351 tkpyfkpgmpyeltvvtvtnp-d--gs-paa-hvp-v-vs-ea--lh-segtt-lsdgt 397
Qy 358 ATPLEKPGIPYKVOVKDSLQVLGCVPIVILNAQTIDVNOETSDLDPSKSVTRVDGV 417
Db 398 akllintpniqslpitvtrnhgdltprqaiksmatatavtqgggsenyhvaiteik 457
Qy 418 ASFVILNPSGVTVLEFNKVTADPLPEENQAREGYRAIAYSSLSQSYLYIDMTDNHALL 477
Db 458 pgdnlpmfnvrgnanslnqikyftylilnkikfykqrrdgnlvtmnlhitpdli 517
Qy 478 VGEHLNI-I-VTPKSPYIDKITHYNYLILSKGIIHFCTREKFSDSASYSQINIPVTQNV 535
Db 518 psfrvayyv-gnn--eivadsvvvdvtdcmgtlvvkgastrddriqkpggaamkile 574
Qy 536 PSSRLVYVYVITGQTAELVSDVWLNIEKCGNQLQVH-LSPDADAY-SPGQTVSLNMA 593
Db 575 gdpqarvgiavdkavvylndkykisqaklwdtiekdfcgtagqgnlgvfedaglal 634
Qy 594 TGMDSWALAVDSAVYGVQCAKKPLERVFQLEKSDLCGCGAGGLNANVFLAGLTF 653
Db 635 ttetnIntkrsaaakcpqpanrrrrsvllldskaskaafqdgqltkccedgmhnpmg 694
Qy 654 LTNANADDSQENDEPCKE-ILRPRT---LQKKIEETAAKYKHSVVRKCCYDGCACVND 709
Db 695 ytekrakyqegdactaaflccchyikgirdenqreselflaredfedelfgddniier 754
Qy 710 -TCQRAARISLSPRCIKAFTECCVWASQLRA-NISHKDMQLCRLHMKTLTPVSRPEI-R 766
Db 755 edfpeawlteeltgpnngiesktvpfyldrdsittwellavgleptkgicvaepyei 814
Qy 767 SYPPESLMEVH-LV--PR-R----KQLQFALPDSLTITWEIQIGISNT-GICVADTVKA 817
Db 815 tvmkdfidrlpysvvkneqveirailynadedi-y-vr---velliynpafcaaste 869
Qy 818 KYFKDVFLEMIPIVSVRGEQILKGTVYNYTSCMQFCVKMSAVEGICTSESPIVHQG 877
Db 870 qy-r-qfpikalsravpfvivypleqglhdveiaavrgelasdgvrkklkvpeger 927
Qy 878 TKSKCVRQKVEGSSRLVTFVLPLEIQLHINFSLETW---FGKEILVKTIRVWPEGVK 935
Db 928 knvrtiieldpskvqg-ggtqeltvianklid-dkvpdtevetriavlgdpvraqiensi 985

Qy 936 RESYSGVTLPD--RGVGTISRKEFPYRIPDLVPKTEIKRILSVKGLVGEILSAVL 993
 Db 986 geklnhiitpogcegmimtpsviaty-yldatogqenlgvdrte-a-ikqimt-g 1041
 Qy 994 QEGINILTHPKGSAAELMSVP-VFYVHYLETGHNHIFHSDDLIEKQKLKKLKEG 1052
 Db 1042 yaqmvvkkadhsyaaftraasawltayvkvlamasmvdkdisheilcggvkvllnr 1101
 Qy 1053 MLSIMSYRNADYSYVWKGSASWLTAFALVLCQVQKVV-EQNQNSICNSLILVENVY 1111
 Db 1102 qpdcqvfknapvhlhgemlqgtkgaep-eas]-tafivtalleesrvckeqlnildssi 1158
 Qy 1112 QLDNGSFKNQSO-QYPIKQGLTDPVEARENSILYTAFTVIGIRKAFDIC-PIVKI-DTAL 1168
 Db 1159 nkatdyllkkyeklqpyttaltayalaaadrlnhdd-r--lm-aa-etgrn--r-w 1207
 Qy 1169 IKADNFULENTLIPAQSTFTLAISAYALSGLDKTHPOF-SIVSALKREALWKNPPIYRFW 1228
 Db 1208 -e--ey-narhn-----iegtayallllmkkkfaevpvrwldidkyvggtyggtq 1257
 Qy 1229 KONLQHKDSSVPNTGARWVETAYALLTSLNKXIDINYVNPVKWLSSEQRVGGGFYSTQ 1288
 Db 1258 atvmvfgaeyeiqmthqdlndisiklperevperysindnavgartvetklnedf 1317
 Qy 1289 DTINAEGLTEYSI-LVKQIRLSMDIDVSKUKALNNYKMDKXFL-GRPVEVLNDDL 1346
 Db 1318 tvsas-gdqkatmtiltvnaqlredanvcnkfhldvsvenvelnkkqkggkaalrki 1376
 Qy 1347 IVSTGFGSLATVHVTVVH-KTSTSEVCS-FYKIDTQDIEASHYRGNSDYKRIVA 1404
 Db 1377 ctry-lg-evds-tm-tiidismaltgfpdaedkrlengvdyriekfeidnmaqgt 1431
 Qy 1405 CASYKPSRESSGSHAVMDISLPTGISANEEDIKALVEGDQLFTDYQI-KD-GH--- 1459
 Db 1432 wviyldkvhaeedclfhkikhfegvfiqpsvkvysynldseqtkfhyhpdketglm 1491
 Qy 1460 VILQMSIPSSDFLCVFRFIELEFVGEFSLPATFVYHRPDQCTWTFYSNLIK--IQ 1517
 Db 1492 kichqncrcaetcsllnqg-k-kidqlriqacagvdyvyktilrieekdgndiy 1549
 Qy 1518 KVCEGAACKVEADCGQMOEELDTISAEYKQKQACKPEIAVAYKVSITSITVENVFKY 1577
 Db 1550 fmdvlevikgt-drnaqakatyvsqrkcealnkldndylinqlsdlwpmk-d-dis 1607
 Qy 1578 KATLLDYKTEGVAEAKDSEITFKVCTNA-ELVKRCRYLING-KEAL-QIKYNSER 1634
 Db 1608 ylit-kn-twierpnedecq-eef-qlncddfaq 1639
 Qy 1635 VYVLDLSLWIEYWPRTTCCSSCQAFIANL-DEFAE 1669

RESULT 4
 ID C03 HUMAN STANDARD; PRT; 1663 AA.
 AC P01024;
 DT 21-JUL-1986 (REL. 01, CREATED)
 DT 21-JUL-1986 (REL. 01, LAST SEQUENCE UPDATE)
 DT 01-OCT-1996 (REL. 34, LAST ANNOTATION UPDATE)
 DE COMPLEMENT C3 PRECURSOR (CONTAINS: C3A ANAPHYLATOXIN).
 GN C3.
 OS HOMO SAPIENS (HUMAN).
 OC EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA;
 OC EUTHERIA; PRIMATES.
 RN (1)

RP SEQUENCE FROM N.A.
 RX MEDLINE; 85140166.
 RA DE BRUIJN M.H.L., FEY G.H.;
 RL PROC. NATL. ACAD. SCI. U.S.A. 82:708-712(1985).
 RN [2]
 RP SEQUENCE OF 672-748.
 RX MEDLINE; 76069169.
 RA HUGLI T.E.;
 RL J. BIOL. CHEM. 250:8293-8301(1975).
 RN [3]
 RP SEQUENCE OF 1409-1563.
 RX MEDLINE; 88154452.
 RA DAUDAKI M.E., BECHERER J.D., LAMBRIS J.D.;
 RL J. IMMUNOL. 140:1577-1580(1988).
 RN [4]
 RP SEQUENCE OF 988-1036.
 RX MEDLINE; 82174534.
 RA THOMAS M.L., JANATOVA J., GRAY W.R., TACK B.F.;
 RL PROC. NATL. ACAD. SCI. U.S.A. 79:1054-1058(1982).
 RN [5]
 RP STRUCTURE BY NMR OF C3A.
 RX MEDLINE; 88276894.
 RA NETTESHEIM D.G., EDALJI R.P., MOLLISON K.W., GREER J., ZUIDERWEG E.R.;
 RL PROC. NATL. ACAD. SCI. U.S.A. 85:5036-5040(1988).
 RN [6]
 RP MUTAGENESIS OF THIOESTER BOND REGION.
 RX MEDLINE; 92250565.
 RA ISAAC L., ISENMAN D.E.;
 RL J. BIOL. CHEM. 267:10062-10069(1992).
 RN [7]
 RP DISULFIDE BONDS.
 RX MEDLINE; 93106233.
 RA DOLMER K., SOTTRUP-JENSEN L.;
 RL FEBS LETT. 315:85-90(1993).
 RN [8]
 RP VARIANT C3F/S.
 RX MEDLINE; 89309808.
 RA POZNANSKY M.C., CLISSOLD P.M., LACHMANN P.J.;
 RL J. IMMUNOL. 143:1254-1258(1989).
 RN [9]
 RP ERRATUM (RETRACTION OF ABOVE ARTICLE).
 RX MEDLINE; 90063087.
 RA POZNANSKY M.C., CLISSOLD P.M., LACHMANN P.J.;
 RL J. IMMUNOL. 143:3860-3862(1989).
 RN [10]
 RP VARIANTS GLY-102 AND PRO-314.
 RX MEDLINE; 91011240.
 RA BOTTO M., YONG FONG K., SO A.K., KOCH C., WALPORT M.J.;
 RL J. EXP. MED. 172:1011-1017(1990).
 RN [11]
 RP VARIANT ASN-549.
 RX MEDLINE; 95050640.
 RA SINGER L., WHITEHEAD W.T., AKAMA H., KATZ Y., FISHELSON Z.,
 RA WETSEL R.A.;
 RL J. BIOL. CHEM. 269:28494-28499(1994).
 RN [12]
 RP VARIANT GIN-1320.
 RA WATANABE Y., MATSUI N., YAN K., NISHIMUKAI H., TOKUNAGA K.,
 RA JUJI T., KOBAYASHI N., KOHSAKA T.;
 RL MOL. IMMUNOL. 30:62-62(1993).
 CC -!- FUNCTION: C3 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE
 CC COMPLEMENT SYSTEM. ITS PROCESSING BY C3 CONVERTASE IS THE CENTRAL
 CC REACTION IN BOTH CLASSICAL AND ALTERNATIVE COMPLEMENT PATHWAYS.
 CC AFTER ACTIVATION C3B CAN BIND COVALENTLY, VIA ITS REACTIVE

CC	THIOLESTER, TO CELL SURFACE CARBOHYDRATES OR IMMUNE AGGREGATES.
CC	-1- SUBUNIT: C3 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 ARG
CC	RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY A DISULFIDE
CC	BOND. C3 CONVERTASE ACTIVATES C3 BY CLEAVING THE ALPHA CHAIN,
CC	RELEASING C3A ANAPHYLATOXIN & GENERATING C3B (BETA CHAIN + ALPHA'
CC	CHAIN).
CC	-1- C3B IS RAPIDLY SPLIT IN TWO POSITIONS BY FACTOR I AND A COFACTOR
CC	TO FORM IC3B (INACTIVATED C3B) AND C3F WHICH IS RELEASED.
CC	-1- IC3B IS THE SLOWLY CLEAVED (POSSIBLY BY FACTOR I) TO FORM C3C AND
CC	C3DG. OTHER PROTEASES PRODUCE OTHER FRAGMENTS SUCH AS C3D OR C3G.
CC	-1- SIMILARITY: TO C4, C5 AND ALPHA-2-MACROGLOBULIN.
CC	-1- FUNCTION: DERIVED FROM PROTEOLYTIC DEGRADATION OF COMPLEMENT C3,
CC	C3A ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS. IT
CC	INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR
CC	PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND
CC	BASOPHILIC LEUKOCYTES.
CC	-1- POLYMORPHISM: THERE ARE TWO ALLELES: C3S (C3 SLOW), THE MOST
CC	COMMON ALLELE IN ALL RACES AND C3F (C3 FAST), RELATIVELY FREQUENT
CC	IN CAUCASIANS, LESS COMMON IN BLACK AMERICAN, EXTREMELY RARE IN
CC	ORIENTALS.
CC	-1- DISEASE: C3 DEFICIENCY CAUSES A SUSCEPTIBILITY TO PYOGENIC
CC	INFECTION.
CC	-1- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN.
DR	EMBL; K02765; G179665; -.
DR	PIR; A01257; C3HU.
DR	PIR; A21603; A27603.
DR	HSSP; P01032; IC5A.
DR	SWISS-2DPAGE; P01024; HUMAN.
DR	MTM; 120700; -.
DR	PROSITE; PS00477; ALPHA 2 MACROGLOBULIN.
KW	COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; PLASMA;
KW	INFLAMMATORY RESPONSE; GLYCOPROTEIN; SIGNAL; POLYMORPHISM;
KW	DISEASE MUTATION.
FT	SIGNAL
FT	1 22
FT	CHAIN 23 1663
FT	CHAIN 23 667
FT	CHAIN 672 1663
FT	PEPTIDE 672 748
FT	CHAIN 749 1663
FT	PEPTIDE 749 954
FT	PEPTIDE 955 1303
FT	PEPTIDE 955 1001
FT	PEPTIDE 1002 1303
FT	PEPTIDE 1304 1320
FT	PEPTIDE 748 749
FT	SITE 954 955
FT	SITE 1303 1304
FT	SITE 1320 1321
FT	DOMAIN 693 728
FT	DOMAIN 1424 1456
FT	DISULFID 559 816
FT	DISULFID 627 662
FT	DISULFID 693 720
FT	DISULFID 694 727
FT	DISULFID 707 728
FT	DISULFID 873 1513
FT	DISULFID 1101 1158
FT	DISULFID 1358 1489
FT	DISULFID 1389 1458
FT	DISULFID 1506 1511
FT	DISULFID 1518 1590
FT	DISULFID 1537 1661
FT	DISULFID 1637 1646
FT	CARBOHYD 85 85
FT	COMPLEMENT C3.
FT	BETA CHAIN.
FT	ALPHA CHAIN.
FT	C3A ANAPHYLATOXIN.
FT	C3B ALPHA' CHAIN.
FT	C3C FRAGMENT.
FT	C3DG FRAGMENT.
FT	C3G FRAGMENT.
FT	C3D FRAGMENT.
FT	C3F FRAGMENT.
FT	CLEAVAGE (BY C3 CONVERTASE).
FT	CLEAVAGE (BY FACTOR I) (POTENTIAL).
FT	CLEAVAGE (BY FACTOR I).
FT	CLEAVAGE (BY FACTOR I).
FT	ANAPHYLATOXIN-LIKE.
FT	PROPERDIN-BINDING.
FT	INTERCHAIN.

[illegible]

Qy 616 AKRQLERFQFLEKSDLGCGAGGJANNANVFLHAGLTLFTLNANADDQENDEPC-KEILR 674
Db 668 rrvavlmeirmdkagdytdgkrlkcedgmrdipmryscqrrarllitqencikafidc 727
Qy 675 PRRTIQ-K-KIEEIAAKYKHSWKKCCYDGAC-VNDETCEQRAARISLPRCIRKAFTEC 731
Db 728 cnhtklreghrdhrlglarsleediipeediisrshfpgswlwtieelkepeknigis 787
Qy 732 CVVASQIRANISHKD-MQJGRHMK-TLIPYSKPEIRBSPESMLWEVH-L-VP-RR--- 783
Db 788 tkvnmiflkdattweilavslkdcgicvadpveirvmqdfidrlrlysvvrneqvei 847
Qy 784 -KQLOPALDLSLTWEIQIGISWT-GICVADTKAVKFDKDFLEMMNIPYSVVRGEQIQL 841
Db 848 ravlfmyteqe-el--kvt-ve-l-l-hnpafcmataknryfqtikippkesvavpyvi 900
Qy 842 KGTVYNYRTSQMQFCVRMSAVEGICTSPVIDHQTKSKCVRQ-KVEGSSHLVFTV 900
Db 901 vplkigqevvkaafnhfisdvkttklvvpegmrinktvaihtldpeklgqgv-qk 959
Qy 901 LPIEIGHNINFSLETW--FKEILVKTLLRVPEVCRESYSGV-TLDPRI-GYTTISRR 956
Db 960 vdv-p-aadlsdqvptdsetril-lqspvvgmaedavdgerlkhllivpagcqeumig 1017
Qy 957 KEFPYRPL-DLVPKTEI-KRLSVKGLLVGEIILSAVISOEGINILTHLPKGSAAELMS 1014
Db 1018 mtpvtviavhldtqewekfgiek-tq-ealel-ikkvtyqlqafkpsayaafnnrpp 1074
Qy 1015 VVPVFFYHLETCGNHNIHSDPLIEKQKUKKLEGMLSIMSTRNADYSYSVWKGCSA 1074
Db 1075 etwltayvvkfslaanliaidshvcagkwllilekqdpdvqcdgpyvhqemiggr 1134
Qy 1075 STWLTAFALRVQVKNKYQNSICNSLLWLVENYOLDNGSEKNSYQPIKLOQTLP 1134
Db 1135 -nakedveltafvllalqeadicqvnslpgslnkageyieasymnlqrpytvaiaq 1193
Qy 1135 VEARENSLYLTAFTVIGIRKAFDIP-LVK-IDTALIKADNFLLNTLPAQSTFTLAISA 1192
Db 1194 yalalmkleepy--lqkfint-a--k-drn--r-w-e-ep-dqql-yn-----veats 1234
Qy 1193 YALSIGDKTHQFRSIVSNLKRREALVKGNPIYRFWKONLQHKDSSVPNTGTARWVETTA 1252
Db 1235 yallallllkdfdvppvrvlneqcyvggystqatfmvfqalagytdvdpdhkdlm 1294
Qy 1253 YALLTSIMLKIDINYNVPVIRKWLSEQRYYGGGYSTQDTINATEGLTEYSLLVQKLR-LSH 1311
Db 1295 dvshlpsrseattfrllwngnllrseetkmeafeltak-gkgrgtlevvayvayhakl 1353
Qy 1312 DIDVSYKUGALHNYKMT-DKNFLGRPEVLELDDLLIVSTGFGSLATVRVTVVHTST 1370
Db 1354 skvtckkdlrlsvlrapetakkpeeaakntfleictky-lq-dvdat-ms--ildismm 1408
Qy 1371 SEEVCS-FYLIKIDTQIEASHYRCYGNSDYKRIVACAS YKPSREESSGSSHAVMDISLP 1429
Db 1409 tgfapdkdllaasgvdrylskymnkafenkntliiylekhteedcltfkvhyfn 1468
Qy 1430 TGISANEEDIKALVEGVDQLTQDI-KD-GH---VILQINSIPSDFLCVRFRIFELFE 1484
Db 1469 vqlipgsvkyvynleesctrfyhpdkddgmleklchsemrcraencf-mqsgoe-k 1526
Qy 1485 VGLSPATFTVYHRPDKQCTMEY-S-TSNIKTQKVEGAACRCVCAOQGMQMEILDUT 1542
Db 1527 inlvrdkacepgvdyvykteltnkllddfdeytmaticqviksgdevqagqqrkfie 1586

Qy 1543 ISAETRRQACKPEIAYAVKVSITSITVENVFVKYKATLLDLYKTCGAVERDSEITFIK 1602
Db 1587 hikrnatklqgkkylmwglas-dl-wgekptseyiigkdtwvehpweaeecqdkyqk 1644
Qy 1603 KVTCTNA-ELVKGROYLKGKALQIKYNSFRYIYPLDSLWIEWPRTTTC-S-SQQA 1659
Db 1645 qceelgaftesmvvygc 1661
Qy 1660 FLANLDEFAEDIFLNGC 1676
RESULT 7
ID CO3 CAVPO STANDARD; PRT; 1666 AA.
AC P12387;
DT 01-OCT-1989 (REL. 12, CREATED)
DT 01-JUN-1994 (REL. 29, LAST SEQUENCE UPDATE)
DT 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE)
DE COMPLEMENT C3 PRECURSOR (CONTAINS: C3A ANAPHYLATOXIN).
GN C3.
OS CAVIA PORCELLUS (GUINEA PIG).
OC EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA;
OC EUTHERIA; RODENTIA.
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE; 90307998.
RA AUERBACH H.S., BURGER R., DODDS A., COLTEN H.R.;
RL J. CLIN. INVEST. 86:96-106(1990).
RN [2]
RP SEQUENCE OF 676-753.
RX MEDLINE; 89113342.
RA GERARD N.P., LIVELY M.O., GERARD C.;
RL PROTEIN SEQ. DATA ANAL. 1:473-478(1988).
RN [3]
RP SEQUENCE OF 993-1032.
RX MEDLINE; 83178889.
RA THOMAS M.L., TACK B.F.;
RL BIOCHEMISTRY 22:942-947(1983).
CC -!- FUNCTION: C3 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE
COMPLEMENT SYSTEM. ITS PROCESSING BY C3 CONVERTASE IS THE CENTRAL
REACTION IN BOTH CLASSICAL AND ALTERNATIVE COMPLEMENT PATHWAYS.
CC AFTER ACTIVATION C3B CAN BIND COVALENTLY, VIA ITS REACTIVE
THIOLESTER, TO CELL SURFACE CARBOHYDRATES OR IMMUNE AGGREGATES.
CC -!- SUBUNIT: C3 PRECURSOR IS FIRST PROCESSED BY THE REMOVAL OF 4 ARG
RESIDUES, FORMING TWO CHAINS, BETA & ALPHA, LINKED BY A DISULFIDE
BOND. C3 CONVERTASE ACTIVATES C3 BY CLEAVING THE ALPHA CHAIN,
CC RELEASING C3A ANAPHYLATOXIN & GENERATING C3B (BETA CHAIN + ALPHA
CHAIN).
CC -!- SIMILARITY: TO C4, C5 AND ALPHA-2-MACROGLOBULIN.
CC -!- FUNCTION: DERIVED FROM PROTEOLYTIC DEGRADATION OF COMPLEMENT C3,
CC C3A ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS. IT
CC INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR
PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND
CC BASOPHILIC LEUKOCYTES.
CC -!- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN.
DR EMBL; M34034; G305335; -.
DR PIR; A37156; A37156.
DR PIR; S03375; S03375.
DR PIR; D20342; D20342.
DR HSSP; P01032; IC5A.
DR PROSITE; PS00477; ALPHA_2_MACROGLOBULIN.
KW COMPLEMENT PATHWAY; COMPLEMENT ALTERNATE PATHWAY; PLASMA;
KW INFLAMMATORY RESPONSE; GLYCOPROTEIN; SIGNAL.
FT SIGNAL 1 29

FT	CHAIN	30	1666	COMPLEMENT C3.
FT	CHAIN	30	671	BETA CHAIN.
FT	CHAIN	676	1666	ALPHA CHAIN.
FT	PEPTIDE	676	753	C3A ANAPHYLATOXIN.
FT	CHAIN	754	1666	C3B (ALPHA' CHAIN).
FT	SITE	753	754	CLEAVAGE (BY C3 CONVERTASE).
FT	DISULFID	557	821	INTERCHAIN (BY SIMILARITY).
FT	DISULFID	630	666	BY SIMILARITY.
FT	DISULFID	630	666	ANAPHYLATOXIN-LIKE.
FT	DOMAIN	698	733	BY SIMILARITY.
FT	DISULFID	698	725	BY SIMILARITY.
FT	DISULFID	699	732	BY SIMILARITY.
FT	DISULFID	712	733	BY SIMILARITY.
FT	DISULFID	878	1517	BY SIMILARITY.
FT	DISULFID	1106	1163	BY SIMILARITY.
FT	DISULFID	1363	1493	BY SIMILARITY.
FT	DISULFID	1394	1462	BY SIMILARITY.
FT	DISULFID	1510	1515	BY SIMILARITY.
FT	DISULFID	1522	1593	BY SIMILARITY.
FT	DISULFID	1540	1664	BY SIMILARITY.
FT	DISULFID	1640	1649	BY SIMILARITY.
FT	THIOLEST	1015	1018	POTENTIAL.
FT	CARBOHYD	944	944	POTENTIAL.
FT	CARBOHYD	1620	1620	POTENTIAL.
FT	CONFLICT	731	731	D -> N (IN REF. 2).
FT	CONFLICT	1013	1013	MISSING (IN REF. 3).
FT	CONFLICT	1018	1018	Q -> E (IN REF. 2).
FT	CONFLICT	1031	1031	MISSING (IN REF. 3).
SQ	SEQUENCE	1666 AA;	186487 MW;	D840333F CRC32;

Db	483	akiryytylimkglkllvgqcpregqalavlmpitkelipfrlvaytylligasaq	542
Qy	495	-KITHYVYLILSKGKI IHFGTRKEFSDASQYSINIDVTQNWPPSSRLIYVYITVGEQTA-	552
Db	543	evwadswadvrdscvgtllvkgqsgkdqdkrqhlprqgmtlriegnqgarvlgvayd	602
Qy	553	ELVSDSWALNTEEKCNQOLQVHLSL--DA-D---ASPCQTVSLNATQMSWALAAND	606
Db	603	kgvfvlnkthkltsqkldwvvekadiqctpgsqdyagvftdaglafskkaglaqtatqe	662
Qy	607	SAYGVQGRGAKEFLERVQFLEKSDLCGCGAGGLNANVFIHLAGLFTLN-ANADDSQEN	665
Db	663	glcdpkaarrtrrsqllmerrmdkagkykselrrccedgmrenpmqfscqrraryvalq	722
Qy	666	DEPC-KEILPRRTLQ-K-KTEETAAKYKHSWKCCYDGCAGVNDE-TCQRAARISUG	721
Db	723	eacvtafidctymaqlrqhrrcqnlgarsdmededlipeediisraqfpefwlwtiee	782
Qy	722	PRCIRAKTECCVWSQLRA-NISHKDMOLGRLLMK-TLLPVSKPEIRSYFPESLWAEVH-	778
Db	783	lkperngistkmmfllkdsaitwellavelsdkkgicvadpfevtvmqdfidrlrpy	842
Qy	779	L-VP-RR---KQLQFALPDSLTLTETIGIGISNT-GICVADTVKARVKDFVLENNIPY	831
Db	843	svrneqweairavlynryrea--q-slkr-vellhnpafcalatakthtqtv---tigpk	896
Qy	832	SVRGEQIOLKCTVNYRTSCMFCVKASVEGICTSESPVIDHQTKSKCRQKVEGS	891
Db	897	ssvavpvyvlpkligqlgevekaavnyfisdgvyktlkvpegmrvnktvairtlnpeq	956
Qy	892	SSHLVTFVPIEIGLHNF--SLTEWFGKEILUKTLRVVPEGVKRESYSGV-TLDPRG	948
Db	957	lgggvy-qreeip-aadlsdgvpdtasetkl-lqgtpvagmaedavdaerklhlitps	1013
Qy	949	I-YGTISRRKEFPYRIP-L-DLVPKTEI-KRILSVKGLVUGELISAVLSQEGINILTHLPK	1005
Db	1014	gcqeqnmgmtptviavhyldtqeufk-f-q---lekrqeaalniinrgyqtqlafkpnwa	1070
Qy	1006	GSAEALMSVVPYVYFHYLETGNHNFISDPLTEKOKLKKLKECMILMSYRNADYS	1065
Db	1071	yaafknrastrwtlyvkvvksfaanligidseivlogavkwlllekqdpdyfqedgpy	1130
Qy	1066	YSWKGGASLTAFALRVLQGVNKVYEQNQNSICNSLLMLVENVOLDNGSKFENSQVQ	1125
Db	1131	hqemiggvrt-agesadveltafvialqeakdicraqvnnleaninkagdyiesryadvr	1189
Qy	1126	PIKLGTLPEVARENSLVITAFVTJGIRKAPDIC-PLVK-IDTALIKADNFLENTLIPAQ	1183
Db	1190	rpytlaiagvalaller-----lng--atlkq-fl-naateknr-weea-rdq---l-yb-	1234
Qy	1184	STFTLAIISAVALSGBKTHPQRSIVSALKREALUKGNPPIYRFWKDNLQHKUSSVPNTG	1243
Db	1235	----veatsyalllaillikdfdvppvrvlneqrygyrgygtqatfmvqfagayqtd	1290
Qy	1244	TARWETTNALLTSJLNUKIDYINVPVKWLSQRYGGGYSTQDTTNALTEGLTYSLL	1303
Db	1291	vpdkhlnmevalqpsrpspskfirlweagsllrseatkneqgfkltak-qkqgqtlev	1349
Qy	1304	VKQLR-LJSDMIDVSYKHKHALHNYKMT-DKNFVIGRPVEVLLINDDLIVSTVGSGGLATVHV	1361
Db	1350	vavyayaktktvcknfdrlvtlkapdptvkkpqaeketmilgictry-lg-dqdat-ms	1406
Qy	1362	TTVWKHSTSEEVG--SFYKIDTQDTEASHRYGVGNISDYKRIIVACASYKPGRSESSGSS	1420

CC C3A ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS. IT

Db	28	ysiitpnvrlleesetfileahdaagdvptvtvqdfll-kkvyitsektvlgatghlnr	86
Qy	23	WISAPKIFRGAZENIVQIYGYTEAFDITISIKSYDPDKFESSYSHVLLSENKTS	82
Db	87	vfkikpaekbefnadkgh-kyrtvvanfsgatvvekaulvafgasyliqrdkiytpgstv	145
Qy	83	ALLTIQP-KQLPGQNPVSYVLEVYLSKHFSKSRMPITYDNGFLIHTDKPVTYDQSV	141
Db	146	fyriftvdmllpvkgtvviwetpdgvpikrdlshagvgiilple-wnipelvmngqv	204
Qy	142	KVRVSYLNDLKPARETVLTFIDPEGSEV--DMVEEIDHIGIISFPDKIPSNRYGMW	199
Db	205	kirafyehapktqfaefevkeyvlpsfevlveptekfy-yihgpkglvsvitarflygk	263
Qy	200	TIKAKYKEDFTTGTAYFENKVEYVLPHPSESIEPEYFNFGYKNF-KNFEITIKARYFNK	258
Db	264	nv-dgtafvifvg-qd--edkkslslslrviiedsgsgeavlrkvlmdgvrpspeal	319
Qy	259	VVTEADVYIFGIREDDKDKQKEMQTAMQNTMLINGTAQVTFDSEYAVKELSYYSLEDL	318
Db	320	vgkelysvetvlihsagsdmvaeersgipitvspyqihfttkpkfkkpampfdlmwvtnp	379

Qy 319 NKYLIAVTVIESGTFSEAEIPGKIVYLSPIKMLVATPLFKGIPYPKIQWQDS 318

Db 380 -d---ge-par-rvp-v-vtqg-ad---aqaltg-ddqvaklsvntnnrqpiltitvstk 426

Qy 379 LQVLVGQVPIVILNAQTIDVNETSNDLPKSVTRVDDGVASEFVILPISGVTLEFNKTD 438

Db 427 kegidparqtatqaqpysthansnyhlsvrvelkpgdnlvnhfhlrtdaqeaki 486

Qy 439 APDLPEENQAREGYRATAYSSLSQSYLYIDWTDNHKA-LVGEHINI-I-VTPKSPYDKI 496

Db 487 rrytylvnmkglklagrvrepqgdvlwlsipitpefiprsfrlvayvtligangrevv 546

Qy 497 THYNLILSKGKLIHFCTREKFSOASQVINPVTQNMWPSRLLYVYVITGEQTA-ELIV 555

Db 547 adsvvvdvdcsvglvlvkgdprdnrpnqapghqtlrlziegnngarvglvadvkgvflnk 606

Qy 556 SDSVWNLINIEKCGNQQLVHLP-DADAYSPQQTVSLNNMATGMSWALAAVDSAVYGVQR 614

Db 607 knltqskidvvekdiaqictpgsknyagvmdagltfktngqlqtdqredpecakpaa 666

Qy 615 GAKXP LERVFOFLEKSDLCGAGGGLNANVHLAGLTELTNANADDSQENDEPC-KEIL 673

Db 667 rrrrsqvlmermdaqytdkglrkccedgmrdlmpyscqrarlitgagcelkafmd 726

Qy 674 RPRRTLQ-K-KTIEIAKYKSHVVKKCCYDGAC-VNDETCQRAARISLUGRCIKAFTE 730

Db 727 cnyitklrqrhrrdvlglarsdvdedlpeediierhfepewltleelkepekngr 786

Qy 731 CCVVASQILRANISHKD-MQGLRLHWK-TLLPVSKPEIRSYFESWLEWH-L-VP-RR-- 783

Db 787 stkmviflksdittweilaveladkqicvadpyeitvmqdfidrlrlypsvvrneqv 846

Qy 784 --KQJQFALPDLTMEIQIGISNT-GICVADTVKAKVFDVLEWNPYPYVVRGSGIQ 840

Db 847 iravlnfyreqek---lkr-vellhnafcmatakkryqti-e-ippkssvavpyvi 900

Qy 841 LKGTVYNYRTSGMFCVKAASAVEGICTSESPVIDHQCKSSKVRQKVGSSSHLVITV 900

Db 901 vlkiglgqvavkaavnfhfiedvqkllkvvpqgmrvnktvavrtldpehlnqvgvqr 960

Qy 901 LPLEIGLHNFLETW--FGKEILVKTLRVPEGVKRESYSGV-TLDPRIY-CTISRR 956

Db 961 dmaadls-dqvdpdsetril-lqgtpvmaedaavdgerlklhltvpsgcqeqnmim 1018

Qy 957 KEFFYPRIPLVPKTEL-KRILSVKGLVGEILSAVLSQEGINILTHLPKGSRAEELMSV 1015

Db 1019 tptviavhlyldtqeqekf-g--lekrqealelikqytqqlafkqplaaayafnnprrps 1075

Qy 1016 VPFYFVYFLETHNHNIFHSDPLIEKQKLRKKLKEGMLSTVSRNADYSYVMKGGAS 1075

Db 1076 twltamwrsfelaanliaidsqvlqavkwliilektpdgvfqedpvhqemiggr- 1134

Qy 1076 TWLTFALFALVLCQNVYEQNSICNSLMLVENVYQLONGSFKENSQYQPIKLQGTLPV 1135

Db 1135 ntkeadvltafvialqeadicqgvmslpgeinkageyleasylnlqrpytvaiagy 1194

Qy 1136 EARENSLYLTAFTYVIGIRKAFDIP-LVK-IDTALIKADNFLENTLPAQSTFLTAISAY 1193

Db 1195 alalmnkleepy--lktkflnt-a--k-drn--r-weepqg----l-yn-----veatsy 1235

Qy 1194 ALSIGDKTHQFORSVSAKREALVGNPPIYRFWKONLQHKDSSVNPVTGARWETAY 1253

Db 1236 alalallikldfsvppvrvlnderyvggygstqatfawfgalagyradvphdklmd 1295

Qy 1254 ALLTSILNKDINYNVPVIRKWLSEDRYGGGYSTQDTINAEGLFEYSLVWKQLR-LSMD 1312

Db 1296 velhlparspvtrfllweagllrseetkqmegfeltak-gkgqgtlsvvtvyhakvkg 1354

Qy 1313 IDVSXKUGALHNYKMT-DKNFLGRPVELLNDLLIVSTGFGSLATVHTVTVVHKTS 1371

Db 1355 ktctckldlrvtkpapatkpkdaksimildicty-lg-dvdat-ms--ildismmt 1409

Qy 1372 EEVCS-FYLIKTDQIDIEASHYRGYGNVDYKRVACASYKPSRESSSGSHAVMDISLPT 1430

Db 1410 gfipdtdllesigvdrylskymdkafsnkntliiykleishaeedclafkvhqfnv 1469

Qy 1431 GTSANEDLALVEGVQLFTDYQI-KD-CH---VILQIASIPSSDFLCVRFRIFELEV 1485

Db 1470 gliqpskvvyeyynleesctrfyhpkkddqmlsKlchnemcrcaenccf-mhqsgd-qv 1527

Qy 1486 GLSPATFTVIEYHRPKQCTMFY-S-TSNIKIQKVGCAKCVADCGQMQEELDUTI 1543

Db 1528 elnerldkacepgvdyvytkklttielsddfdeytmteqviksgsdevqagqerrfish 1587

Qy 1544 SAETRKQTAKEFIAYAYKVSITSITVENFVKYKATLLDIYKTGEAVAEKDEITFIKK 1603

Db 1588 vkcnalklqkqkylmwglss-dl-wgekptsyigkdtwvehwp 1632

Qy 1604 VTCTNA-ELVKGRQYLIMGKEALQIKYNEFSFRYIYPLDSLTLTWIEYWP 1649

RESULT 9

ID CO3_EPTBU STANDARD; PRT; 1620 RA.

AC P98094;

DT 01-NOV-1995 (REL. 32, CREATED)

DT 01-NOV-1995 (REL. 32, LAST SEQUENCE UPDATE)

DT 01-FEB-1996 (REL. 33, LAST ANNOTATION UPDATE)

DE COMPLEMENT C3 (CONTAINS: C3A ANAPHYLATOXIN (FRAGMENT)).

GN C3.

OS EUPHRETUS BURGERI (INSHORE HAGFISH).

OC EUPHRYOTA; METAZOA; CHORDATA; VERTEBRATA; PISCES;

OC AGNATHA (CYCLOSTOMATA).

RN [1]

RP SEQUENCE FROM N.A., AND PARTIAL SEQUENCE.

RX MEDLINE; 92192016.

RA ISHIGURO H., KOBAYASHI K., SUZUKI M., TITANI K., TOMONAGA S., KUROSAWA Y.;

RL EMBO J. 11:829-837(1992).

CC -!- FUNCTION: C3 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE COMPLEMENT SYSTEM. AFTER ACTIVATION (C3B), IT CAN BIND COVALENTLY, VIA ITS REACTIVE THIOLESTER, TO CELL SURFACE CARBOHYDRATES OR IMMUNE AGGREGATES. CYCLOSTOMATES C3 APPEARS TO REPRESENT THE COMMON ANCESTOR OF MAMMALIAN C3 AND C4, SHOWING SIMILARITIES TO BOTH PROTEINS.

CC -!- SIMILARITY: TO C4, C5 AND ALPHA-2-MACROGLOBULIN.

CC -!- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN.

DR EMBL; 211595; G62775; -.

DR EMBL; 211596; -; NOT ANNOTATED_CDS.

DR PIR; S21045; S21045.

DR PROSITE; PS00477; ALPHA 2 MACROGLOBULIN.

KW COMPLEMENT PATHWAY; PLASMA; INFLAMMATORY RESPONSE; GLYCOPROTEIN; SIGNAL.

KW NON TER 1 1

FT CHAIN <1 1620 COMPLEMENT C3.

FT CHAIN <1 633 BETA CHAIN (BY SIMILARITY).

FT CHAIN 634 1336 ALPHA CHAIN (BY SIMILARITY).

FT CHAIN 1343 1620 GAMMA CHAIN (BY SIMILARITY).

FT PEPTIDE 634 714 C3A ANAPHYLATOXIN (BY SIMILARITY).

Qy 831 YSVRGQIQGLQVTVYNTSCMOFCVKMSAVEGICTSESPVIDHQGTSKCVQRKVEG 890
 Db 871 mstrvpyviipmklglhsievksavknsgndgvrklrrvaeqvlvkketnvlhnp-v 929
 Qy 891 SSSHLVTVFVLPJLGIJHNF--SLETFKGKELUWTLVVPWEGKRGESYGVTLDPRG 948
 Db 930 khgg-eqthlpsvprnqpsnadsdlstvtagevtqlveqalsdglslvqpvqc 988
 Qy 949 IYGTISRREPPYIPIDIVPKTEKILSVK-GLLVGEILSNVLSQECINILTHLPKGS 1007
 Db 989 geqmiamtlypviat-hyldntkkwedi-q--ldkrntalkviniqyqlayrkedqsy 1044
 Qy 1008 AEAEIMSV-VPVYVHYLETGNHNIIFSDPLIEKOKIKKIKKEGMISIMSRADYSY 1066
 Db 1045 aawsvrqsatltayvvkvfamsstllisqenvlctavkwlilntqpqdgifnefapvih 1104
 Qy 1067 SVMRGGSASTWLTAFALRVLCQVKNKYVEQNQNSICNSLLWLVENYQLDNGSFKENSQYOP 1126
 Db 1105 aemtgn--vrgedndaamtafvliamqeaasvceqsvslpagsmakavayl-ekrlp-h- 1159
 Qy 1127 IKLQGTLPVEARENSLYLFTFVIGIRKAFDICPL-VK-IDTALLKADNLENTLETPAQ5 1184
 Db 1160 ---lt-npyavam--taya-l--anaqlnkletllkfasp-----q-lbh--wvpg-gy 1201
 Qy 1185 FTIATISATSLSGDTHQFQFISVSLAKREALVKNPPIYRFHFKDNLQHKUSSVPNTGT 1244
 Db 1202 qytleatsvallalvkkafeaaqpvrvlnkqkvqvgystqetlmvfqavaeywshv 1261
 Qy 1245 ARMYETTAYALTSLANKDINTVNPVKMLSEQRGGGYSTQDTTINATELGEYSLLV 1304
 Db 1262 kllkdfdmninlvagrasvtnnknqfhttrtdkvnssdkdltkvae-gngeatlsv 1320
 Qy 1305 KQLR-LSMDIDVSYKHKGALHNYKMTDKN-FLGRPVEVL-LNDDLVSTGSGSLATVHV 1361
 Db 1321 vlyyalpeeksdcesfdlvtltkmdktshd-akesfmltie-vlykns--orda-t 1375
 Qy 1362 TTVVHTSTSEEV-C-SFYLIKIDTDIEASHYRGVGNSDYKRIVACASYKPSRESSSGS 1419
 Db 1376 m-sildigltgftvtdddlnqlekgtryekfemdkvlsgerglilyldkvskhledr 1434
 Qy 1420 SHAVNDISLPTGISANEEDLKALVEGVQDLFTDYQI-K--D-GHVLQINSIPSSDFLC 1474
 Db 1435 lsfkhrvqvglqpaavsvyeyn-qkrvkvfhpqreggtlarlclgdvctcaesc 1493
 Qy 1475 VRFRIEFLFVGLSPATTVYEHPRDKQCTMFY-STSN1-KIQKCEGAACKCVCVAD 1532
 Db 1494 s-mktgepdv--q-ridkaqagldvyvkatvdfsklthtdtytkidvi-kpqtde 1548
 Qy 1533 QMOQEELDTISAEKTRQACKPEIAYVAKVSIITSITVENVEVYKATL-LDYITGEAV 1591
 Db 1549 gveqndrfmglaycrealglnqkqymimgksedlhvrvedkllgqykvlgqetwieww 1608
 Qy 1592 AEKDESLTEIKKVCTNA-ELVKGROVLINGK-EALQ-IKYNFSFRYIPDLISLWIEYW 1648
 Db 1609 pseqcterdvrevclgidefinqitfqc 1638
 Qy 1649 PRDTTCSS--COAFLANLDEAFEDIFLNGC 1676

RESULT 11
 ID C04 HUMAN STANDARD; PRT; 1741 AA.
 AC P01028;
 DT 21-JUL-1986 (REL. 01, CREATED)

DT 01-NOV-1990 (REL. 16, LAST SEQUENCE UPDATE)
 DT 01-OCT-1996 (REL. 34, LAST ANNOTATION UPDATE)
 DE COMPLEMENT C4 PRECURSOR (CONTAINS: C4A ANAPHYLATOXIN).
 GN C4A AND C4B.
 OS HOMO SAPIENS (HUMAN).
 OC EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA;
 OC EUETHERIA; PRIMATES.
 RN [1]
 RP SEQUENCE OF 1-22 AND 1056-1225 FROM N.A.
 RX MEDLINE; 85156269.
 RA BELT K.T., YU C.Y., CARROLL M.C., PORTER R.R.;
 RL IMMUNOGENETICS 21:173-180(1985).
 RN [2]
 RP SEQUENCE OF 20-1741 FROM N.A.
 RX TISSUE=LIVER;
 RA BELT K.T., CARROLL M.C., PORTER R.R.;
 RL CELL 36:907-914(1984).
 RN [3]
 RP SEQUENCE OF 680-756.
 RX MEDLINE; 81264286.
 RA MOON K.E., GORSKI J.P., HUGLI T.E.;
 RL J. BIOL. CHEM. 256:8685-8692(1981).
 RN [4]
 RP SEQUENCE OF 957-1044.
 RX MEDLINE; 82182029.
 RA CAMPBELL R.D., GAGNON J., PORTER R.R.;
 RL BIOCHEM. J. 199:359-370(1981).
 RN [5]
 RP SEQUENCE OF 990-1037.
 RX MEDLINE; 82150875.
 RA HARRISON R.A., THOMAS M.L., TACK B.F.;
 RL PROC. NATL. ACAD. SCI. U.S.A. 78:7388-7392(1981).
 RN [6]
 RP STRUCTURAL BASIS OF POLYMORPHISM.
 RX MEDLINE; 87080272.
 RA YU C.Y., BELT K.T., GILES C.M., CAMPBELL R.D., PORTER R.R.;
 RL EMBO J. 5:2873-2881(1986).
 RN [7]
 RP VARIANT C4A6 ALLOTYPE.
 RX MEDLINE; 92242905.
 RA ANDERSON M.J., MILNER C.M., COTTON G.H., CAMPBELL R.D.;
 RL J. IMMUNOL. 148:2795-2802(1992).
 CC -!- FUNCTION: C4 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE
 CLASSICAL PATHWAY OF THE COMPLEMENT SYSTEM. IT IS PROCESSED BY
 ACTIVATED C1 WHICH REMOVE FROM THE ALPHA CHAIN THE C4A
 ANAPHYLATOXIN.
 CC -!- SUBUNIT: THIS PROTEIN IS SYNTHESIZED AS A SINGLE-CHAIN PRECURSOR
 AND, PRIOR TO SECRETION, IS ENZYMATICALLY CLEAVED TO FORM A TRIMER
 OF NONIDENTICAL CHAINS (ALPHA, BETA, AND GAMMA).
 CC -!- POLYMORPHISM: HUMAN COMPLEMENT COMPONENT C4 IS POLYMORPHIC WITH AT
 LEAST TWO LOCI, C4A & C4B. 13 ALLELES OF C4A & 22 ALLELES OF C4B
 HAVE BEEN DETECTED. THE ALLELE SHOWN HERE IS C4A4.
 CC -!- C4A ALLOTYPES REACT MORE RAPIDLY WITH THE AMINO GROUP OF PEPTIDE
 ANTIGENS WHILE C4B ALLOTYPES REACT MORE RAPIDLY WITH THE HYDROXYL
 GROUP OF CARBOHYDRATE ANTIGENS.
 CC -!- POLYMORPHISM: THE C4A ALLELES CARRY THE BLOOD GROUP RODGERS WHILE
 THE C4B ALLELES CARRY THE BLOOD GROUP CHIDO.
 CC -!- DISEASE: THE C4A6 ALLOTYPE IS TOTALLY DEFICIENT IN HEMOLYTIC
 ACTIVITY.
 CC -!- FUNCTION: DERIVED FROM PROTEOLYTIC DEGRADATION OF COMPLEMENT C4,
 C4A ANAPHYLATOXIN IS A MEDIATOR OF LOCAL INFLAMMATORY PROCESS. IT
 INDUCES THE CONTRACTION OF SMOOTH MUSCLE, INCREASES VASCULAR
 PERMEABILITY AND CAUSES HISTAMINE RELEASE FROM MAST CELLS AND

Qy 1231 NIQKD-S-S-VP-N-TGT---AR-M-VETIYALLTSL-NIKDINTVNPVIMLSEQR 1279
Db 1295 fggfstcdvialdaleaywiashteerglnvltsetgrngfksahqlqnnrqirgl 1354
Qy 1280 YGGGFYSTODTINAIEGLTYSLL--VKQIR-LSMDID-VS---YK-HKCALFN--YK-M 1328
Db 1355 eelqfslgskinvkvngskgtikvlrtnvldmktctqdlqievtkghvymean 1414
Qy 1329 TDK-NF-LGRPVLL--ND-D-LIVSTGGS-GL--AT---VHV-TTV-VHKTSTEEV 1374
Db 1415 edveydelpakddpqlqplqfegrrrrrreapkvveeqesrvhytvcvwrngk 1474
Qy 1375 CSF-Y-L-KIDTQD-IEA-SHRGY-GNSDYKR-----IVA---C-ASYKPS-REES 1416
Db 1475 vqlgmaiadvtllqfghalradlektsldryshfeteqhvllfydsvpter-ecv 1533
Qy 1417 SG-SSHAVNDISLPTGISANEEDLKALVEGVQDLFTDYQIKDGHVILQINSIPSSDFLCV 1475
Db 1534 gfeavqevpvlvpasatlvdynperrcsfvgapskerlllatlcsaevcqaegtcv 1593
Qy 1476 REFIFELFEVGLSPATFTVVEYHRPDKQCTMFYST-SNIKI-QKVCEGAACKVEADCG 1533
Db 1594 tqralergldedgymkfacyprvevfgvkvredraafrlfetkitcvlhfktk 1653
Qy 1534 QMQEELDTI-SAET-RKQACK-PEIAYAVKVSITSITVENVFYKATLDDIYKTGEA 1590
Db 1654 vkaanqmrnfivraec-frllepkeylimldg--atvdlqghpqyldlenswiepm 1710
Qy 1591 VAEKDSEI-TFKKVTCTNAELVKRGYLLMGKCALQKYNESFRYIYPLDSLTWIEWP 1649
Db 1711 eerlcrstrqaacqndflqeygtgc 1739
Qy 1650 RDTTC-SSCQ-AFLANLDEAFIDFNGC 1676
RESULT 12
ID C04 MOUSE STANDARD; PRT; 1738 AA.
AC P01029;
DT 21-JUL-1986 (REL. 01, CREATED)
DT 01-AUG-1988 (REL. 08, LAST SEQUENCE UPDATE)
DT 01-OCT-1996 (REL. 34, LAST ANNOTATION UPDATE)
DE COMPLEMENT C4 PRECURSOR (CONTAINS: C4A ANAPHYLATOXIN).
GN C4.
OS MUS MUSCULUS (MOUSE).
OC EUKARYOTA; METAZOA; CHORDATA; VERTEBRATA; TETRAPODA; MAMMALIA;
OC EUTHERIA; RODENTIA.
RN [1]
RP SEQUENCE FROM N.A.
RX MEDLINE; 85298264.
RA SEPICH D.S., NOONAN D.J., OGATA R.T.;
RL PROC. NATL. ACAD. SCI. U.S.A. 82:5895-5899(1985).
RN [2]
RP SEQUENCE FROM N.A.
RC STRAIN=B12.WR;
RX MEDLINE; 87309760.
RA ROSA P.A., SEPICH D.S., ROBINS D.M., OGATA R.T.;
RL J. IMMUNOL. 139:1568-1577(1987).
RN [3]
RP SEQUENCE FROM N.A.
RC STRAIN=B12.WR; TISSUE=LIVER;
RX MEDLINE; 89380278.
RA OGATA R.T., ROSA P.A., ZEPF N.E.;
RL J. BIOL. CHEM. 264:16565-16572(1989).
RN [4]

RP SEQUENCE FROM N.A.
RC STRAIN=FW; TISSUE=LIVER;
RX MEDLINE; 85289294.
RA NONAKA M., NAKAYAMA K., YEUL Y.D., TAKAHASHI M.;
RL J. BIOL. CHEM. 260:10936-10943(1985).
RN [5]
RP SEQUENCE OF 651-810 AND 924-1083 FROM N.A.
RX MEDLINE; 85038607.
RA NONAKA M., TAKAHASHI M., NATSUUME-SAKAI S., NONAKA M., TANAKA S.,
RA SHIMIZU A., HONJO T.;
RL PROC. NATL. ACAD. SCI. U.S.A. 81:6822-6826(1984).
RN [6]
RP SEQUENCE OF 1105-1449 FROM N.A.
RX MEDLINE; 85166208.
RA LEVI-STRAUSS M., TOSI M., STEINMETZ M., KLEIN J., MEO T.;
RL PROC. NATL. ACAD. SCI. U.S.A. 82:1746-1750(1985).
RN [7]
RP SEQUENCE OF 1257-1376 FROM N.A.
RX MEDLINE; 85038859.
RA TOSI M., LEVI-STRAUSS M., DUFONCHEL C., MEO T.;
RL PHILOS. TRANS. R. SOC. LOND., B, BIOL. SCI. 306:389-394(1984).
RN [8]
RP SEQUENCE OF 1360-1511 FROM N.A.
RX MEDLINE; 83273751.
RA OGATA R.T., SHREFFLER D.C., SEPICH D.S., LILLY S.P.;
RL PROC. NATL. ACAD. SCI. U.S.A. 80:5061-5065(1983).
RN [9]
RP SEQUENCE OF 1-128 FROM N.A.
RC STRAIN=FW; TISSUE=LIVER;
RX MEDLINE; 86031969.
RA NONAKA M., NAKAYAMA K., YEUL Y.D., SHIMIZU A., TAKAHASHI M.;
RL IMMUNOL. REV. 87:81-99(1985).
CC -!- FUNCTION: C4 PLAYS A CENTRAL ROLE IN THE ACTIVATION OF THE
CC CLASSICAL PATHWAY OF THE COMPLEMENT SYSTEM. IT IS PROCESSED BY
CC ACTIVATED C1 WHICH REMOVE FROM THE ALPHA CHAIN THE C4A
CC ANAPHYLATOXIN.
CC -!- SUBUNIT: THIS PROTEIN IS SYNTHESIZED AS A SINGLE-CHAIN PRECURSOR
CC AND, PRIOR TO SECRETION, IS ENZYMATICALLY CLEAVED TO FORM A TRIMER
CC OF NONIDENTICAL CHAINS (ALPHA, BETA, AND GAMMA).
CC -!- C4 IS A MAJOR HISTOCOMPATIBILITY COMPLEX CLASS-III PROTEIN.
CC -!- SIMILARITY: TO C3, C5 AND ALPHA-2-MACROGLOBULIN.
CC -!- SIMILARITY: CONTAINS ONE ANAPHYLATOXIN-LIKE DOMAIN.
DR EMBL; K00019; E19386; -.
DR EMBL; M11729; G387420; -.
DR EMBL; M12968; G199270; -.
DR EMBL; M12970; G199262; -.
DR EMBL; M12972; G199264; -.
DR EMBL; M11789; G387438; -.
DR EMBL; K02798; G199282; -.
DR EMBL; M17440; G387440; -.
DR PIR; A01264; A01264.
DR PIR; A21692; A21692.
DR PIR; A20339; A22039.
DR PIR; A24558; A24558.
DR PIR; A29059; A29059.
DR HSSP; P01032; 1C5A.
DR PROSITE; PS00477; ALPHA 2 MACROGLOBULIN.
KW COMPLEMENT PATHWAY; PLASMA; GLYCOPROTEIN; MHC III; SIGNAL;
RN INFLAMMATORY RESPONSE.
FT SIGNAL 1 19
FT CHAIN 20 673 BETA CHAIN.
FT CHAIN 678 1443 ALPHA CHAIN.
FT CHAIN 1448 1738 GAMMA CHAIN.
FT PEPTIDE 678 753 C4A ANAPHYLATOXIN.


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Db 67 ivtsgsgceqnmiamhtviahvlyldhteqwdf-s--lekqaelelikgytqqlafk 123
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RA FEY G.H.;

RA FEY G.H.

	1131	lesswmiieggnfgyvfkalmayafalgagnqekr--neilksldkea-ikednsihwer	1188
Db		: : :	:
	1169	IRADNFULENTLPAQSTFTTIAISAYALSIGDKTHQFRSIVSALKREALKGNPPITYRWF	1228
Qy		: : :	:
	1189	pqkptkseegylytpqasaeavemsyvvarlktagpaspedsalemgtkwltkqmqsy	1248
Db		: : :	:
	1229	KDNLQHKKDSV--PNTGTARWETAYAJ--LTSL---NKLDINYVPVKWLSEQR-Y	1280
Qy		: : :	:
	1249	gg-festqdtvwaldalekyga-atfsksqtpsvtvgsgsfqskfvdkenzlllqqv	1306
Db		: : :	:
	1281	GGGFYSTQDTINAIIEGLTEYSLVLKQLRLSDMDIDVSZYKKGAL-HNYKMTDN-FLCRPV	1338
Qy		: : :	:
	1307	slypignytvs=-gegcyyaq-ttlcynvplekqpafkalvtqvplctnmpkg-qms	1363
Db		: : :	:
	1339	EV-LUNDDLVITGFGSGLAETHVTHTTTSTSEEVCFSYLIKIDTQDIASHYRTGNS	1397
Qy		: : :	:
	1364	-fqiele-isymgst--pas-nm-viadvkmlsgfilpktvvkl--erlighv-srtevt	1415
Db		: : :	:
	1398	DYKRIVACASYKPREESSGSHAVMDISLPTGTISANEEDJAKALVEGDQDLFTDQIKD	1457
Qy		: : :	:
	1416	nrvillyldqv-tnqtlsfiiqqdipkylqpaiukvydyetd	1459
Db		: : :	:
	1458	GHVILQNSIPSSDFLCVRERIFELFEVGLSPAFFTYVEYHRPD	1502
Qy		: : :	:

[illegible]

Query Match	5.5%;	Score 661;	DB 1;	Length 1476;
Best Local Similarity	24.1%;	Pred. No. 3.83e-116;		
Matches	368;	Conservative 407;	Mismatches 612;	Indels 140;
Gaps	112;			
Db	28	dskymvlvpsqlytpekiicllhlyqndntvttaelvsqegrknlfde-lv-lakdl-f 84		
Qy	20	EQTWVISAPKIFRVGASENIWIOVGYCTAFDATISIKSYDPDKFSYSSCHVHLSSENKF 79		
Db	85	qcvsfi-l-pr-lssdee-dflvydkgpthefskrkavlnknesvfvctdkpykvp 140		
Qy	80	ONSALTITQPKQLPGQGNPSVYVLEV-VSKH-FSKSKRMPITYDNGFLFIHTDKPVYTP 137		
Db	141	gqsckvrvmgdmkmlrplnellpalyedpkknri-mqwrldiktenglkqmsfslsaepi 199		
Qy	138	DQSKVRVYSLINDLKPAKRETWLF1-DPGCESEVMV-EEDIHGIIISFPDFKIPSNPR 195		
Db	200	qgpykiwv-hkesgekeehs-ftvmefvlprfnvdlkvp-namsvmd-evlstaacqkyt 255		
Qy	196	YGMWIRAKYKEDFSTTGATAYFEWKEYVLPHFVSVEPEYNIYGNKFNKFEITIKARYF 255		
Db	256	yqkypv-ghvkinv-cret-etgcrev-nsqldnng-cs-tqevni-telqskk-rnyev 307		
Qy	256	YKRWTEADVYITGIRREDLKDDQKEMQWQTAQNTMLINGIAQTWTFDSEITAKELSYSL 315		
Db	308	q-l---f-hvnatvteegtql-efarsgttkieritnklifkadehf-rhgiptfvkvr 360		
Qy	316	EDLNKNKYDYTAVTVIESTGCGSEAEIFGIYVUSPYKUN-LVATP1FLKPGIPIPKVQ 374		
Db	361	lvdiqgdiponekfikaqelsytaattdhghlaefsidtcsagsslhikvnhkeeds 420		
Qy	375	VRKSDLDQVGGVPEVILNAQTIDVNOETSDLDPSKSVTRVDVG-VASEFVILNPISGVTVLEF 433		
Db	421	csyfyemeethasaakhvay-av-yg-leksyilyld-tetsailpcnqihvtqahfllkgd 476		
Qy	434	NVKTDAPDLPFENQAREGYRAIAYSLSQSYLYIDWTDNHKALLVGE-H-1NIIVTPKSP 491		
Db	477	lgvlelfiylvmaagsgiiqctgnthqvegeapvkqfaleipvesvmymakmliyt 536		
Qy	492	Y-IDK1TH7NYLLISKGK1HFG--TRE-KFSDAQY--SIN1PTQNNVPSRLIAYY 544		

FT	DISULFID	48	86	BY SIMILARITY.
FT	DISULFID	251	276	BY SIMILARITY.
FT	DISULFID	269	288	BY SIMILARITY.
FT	DISULFID	461	555	BY SIMILARITY.
FT	DISULFID	587	773	BY SIMILARITY.
FT	DISULFID	634	680	BY SIMILARITY.
FT	DISULFID	849	885	BY SIMILARITY.
FT	DISULFID	923	1323	BY SIMILARITY.
FT	DISULFID	1081	1129	BY SIMILARITY.
FT	DISULFID	1354	1469	BY SIMILARITY.
FT	THIOLEST	974	977	BY SIMILARITY.
FT	CARBOHYD	55	55	POTENTIAL.
FT	CARBOHYD	294	294	POTENTIAL.
FT	CARBOHYD	313	313	POTENTIAL.
FT	CARBOHYD	500	500	POTENTIAL.
FT	CARBOHYD	749	749	POTENTIAL.
FT	CARBOHYD	776	776	POTENTIAL.
FT	CARBOHYD	871	871	POTENTIAL.
FT	CARBOHYD	993	993	POTENTIAL.
FT	CARBOHYD	1142	1142	POTENTIAL.
FT	CARBOHYD	1180	1180	POTENTIAL.
FT	CARBOHYD	1426	1426	POTENTIAL.
SQ	SEQUENCE	1476 AA;	165139 MW; 7E3EAF00 CRC32;	

Key	Location/Qualifiers
Peptide	1..22
/note= "Signal sequence"	
Protein	23..655
/note= "Alpha-chain"	
Modified site	153
/note= "Potential glycosylation site"	
Modified site	158
/note= "Potential glycosylation site"	
Modified site	210
/note= "Potential glycosylation site"	
Protein	739..1269
/note= "Gamma-chain"	
Binding site	742..751
/note= "Factor B binding site"	
Active site	996..1017
/note= "Thioester site"	
Binding site	1176..1201
/note= "Factor H binding site"	
Binding site	1214..1219
/note= "Potential CR2 binding site"	
Binding site	1221..1235
/note= "Factor H binding site"	
Protein	1270..1651
/note= "Beta-chain"	
Modified site	1352
/note= "Potential glycosylation site"	
Binding site	1419..1442
/note= "Properdin binding site"	
PN	W09423024-A.
PD	13-OCT-1994.
PF	07-APR-1994; U03441.
PR	07-APR-1993; US-043747.
PI	(GEU) UNIV GEORGETOWN.
PI	Breddehorst R, Fritzing DC, Vogel

Db	4	malylvaa1ligfpg-s-shaalyt litpavl rtdteeqilv eahgdstpk sldifvndf	61
Qy	1	MGLLG1ICFLI-FLGKTWCQEQVTYISAKIFRVCASENIVIQVYGYTEAFDATISIKSY	59
Db	62	prkqktlitfgsvdmnmgasfmvtpl-kypake lnkds kmgvkvvkvgtgqvalekvvl	120
Qy	60	PDKKSYS5SGUHLIS-SENKQNSA LITTPQKIPQCGNPVSVYILEVSKHPSKSRMP	118
Db	121	lsyqsgvfictdkgiytpgspvryrvfsvdnhmhmrmdktviefqtpegivvs-ekpvn	179
Qy	119	ITYDNGLFIHTDKPYPTDQSVKRVRYSLNDLKP AKRETVLITFIDPEGSEYDMVEEID	178
Db	180	psgair-p-ynl pelvsfgtkavakvshs peesytay fdrveylvpsfevrlqpedkfl	237
Qy	179	HIGISFPDFKIPSNRGMWTIKAKYKEDFSTGCTTAYFEWKEYVLPHFVSIEPEYFI	238
Db	238	-yidgnknfhsitarylygkkv-egvafvfgvk--l-ddakkeipds ltriplidgdg	290
Qy	239	GYN-KFNFEITIKARYFNKVTYADVITFGREDLDDKQKEMQTAQNMTLINGIA	297
Db	293	eatlkrdt-lrs-rfdlnqlvghtlysvtvtitesgdmvvtcggihvtpespylyft	350
Qy	298	QVTFDSEYAKVLSYSLIEDLNKKYLYIAVTVIESTGGFSEAEIPGIKVLYSPYKMLIV	357
Db	351	ktkyfkgmpyelvtvytnp-d--gs-paa-hvp-v-vs-ea--lh-segtt-lsdgt	397
Qy	358	ATPLFKGIPYPIKVQKDSLQLVGGVPVILNAQTI DVNQETSLDLPDPSKSVTRVDDGV	417
Db	398	akliintplnqlspitvrthngdlprerqaksmatayqtqggseny lhwaitetcik	457
Qy	418	ASVNLPLSGVTVLEFNKVTADPLPEENQARECYRAIAYSSLSQSYLYIDMTDNHALL	477
Db	458	pqdnlpvnfnvrngnanslnqikyftylilnkakikfkvqrqprirdqnlvcmllhtpdli	517

Qy	478	VGEHLNI-I-VTPKSPYDKITHYNYLILSRKGIHFGTRKREXSDASYQSINIPYTONWV	535
Db	518	psfrfvyayqy-gnn--eivadvsvvdkvdkcmgtllvvgaesrddridhtkpaamkile	574
Qy	536	psrlllyvvtgeotaelvsvlwnieekgnqlqvh--lspdaday-spgqvtvsnma	593
Db	575	gdbqrgvlvadvkavylndkykiegaklwdiekadfgctagsqnnlgvfedaglal	634
Qy	594	TCMDSWALAAVDASYVGYQVCAKPKLERVFQFLKSDJLCCGAGGGLNNANVFLAGLTF	653
Db	635	tstetnltkqrsaakcpqpnrrrsrvsllldskasakaafgdqglrkccedgmhndpmg	694
Qy	654	LTVANADSQZNDPECKE--ILRPRT---LQKKITEEIAKYKHSVVKKCYDGACVWNDE	709
Db	695	ytcekrakyiqegackaaflecchkykigidenqreselflaredfedelfgdndliar	754
Qy	710	--TCQRAARISLPGRCIKAFTECCVWASQRA--NISHKOMLGRLLHMKTLPLVPKPEI-R	766
Db	755	adpseawltteeltgepmpngisestvpyfyrldsttwellawlgloptkgicvaepvei	814
Qy	767	SYTPESLWEVH-LV--PR-R---KQLQFALPDSLTLWEIQIGISNT--GICVADTVKA	817
Db	815	vmkdfidldrlpysvvnkeqveiralynvadedi-y-vr---velinyapfaeateg	869
Qy	818	KVFKDVFLENNIPYSVVRGEQIQLGTVNYRTSGMFCVKMSAVEGICTSESPVIDHQG	877
Db	870	qiy-r-qgfipkalsarapfviplqegldhveiasvrgeslaedgvrrklkvvpeger	927
Qy	878	TKSKCKQVKVEGSSHLVTVLPLEIGLNNFSLFW--FKGEILWKTLLVWPGGVK	935
Db	928	knivtielldpsvkvy-ggtqeltvianklid-kvnpdtevetriavlgdpvqaaiensid	985
Qy	936	RESYSGVTLDP--RGYGTITSRKKEFYRIPIDLPVTPKTRILSVKGLVAGEILSAVIS	993
Db	986	gsklnhliitpscgcegmmtcpsviaty-yldatqwenlgvdrtte-a-ikqimt-q	1041
Qy	994	QEGINILPHLPKGSAAELMSVWP-VFYVHYLETGNHNNIFHSDPLIEKQKIKKKLKEG	1055
Db	1042	yaqgmvykkadhyaafntnrassawltayvkvvlamasnmvkdshheilcgvgkwilnr	1101
Qy	1053	MLSINSYRNADYSYVMKSGSASTLTAFALRVLGVQNKVY--EQNQNSICNSLLWVNY	1111
Db	1102	qgpdgvfkenapvhiqemlggtkgaep-easl--tafi tallaesrsvckeginlidsai	1158
Qy	1112	QLDNGSKFENSQ-YQP IKLGCTLPVARENSLYTAFVIGIRKAFDIC-PLWKI-DTAL	1168
Db	1159	nkadyllkkyeklqrpyttaltayalaaadlndd-r-v-v--lm-aa-stgrn---r-w	1201
Qy	1169	IKADNFLENTLPAOSTFTTLTAISAYALSIGKTHQPFQFRISVALKREALVKNPPIYRFW	1228
Db	1208	e--ey-narthn-----tegteyalallkmkkfaevgppvrvllidkkyvggtvgqtc	1257
Qy	1229	KDNLQHKDSSVPNTGTARMVETTAYALLTSLNLKDINYVNPVKWLSEQRYYGGCFSTQ	1288
Db	1258	atvmvfgaeyeiqmthqdnldisiklperrepvrysindrnvqartvektlnefd	1317
Qy	1289	DTINAIEGLTEYSI--LVKQILRLSMDIDVSYKKHGLAHNYKMTDKNFJ--GRPVEVLLNDL	1346
Db	1318	tlvaas-gdgkatmtiltvynaqldredanvcnkhldvsvenvelnlkqagkgaaalrki	1376
Qy	1347	IVSTGFGSGLATVHTTVVH-KYTSSTEEVCS-FYIKIDQTDQIEASHVRVCGNSDYKEIVA	1404
Db	1377	ctiv-lq--evds-tm-tiidiqsmktgffdoaedlkrilenavdriviekfeidnmackot	1431

	I : I : : : :::: I : :: : : :: :: :: ::	
Qy	1405 CASKYPSREESSGSSHAVMDISLPTCISANEEDIKALVEGVQDLFTDYQI-KD-GH----	1459
Db	1432 vviylkdwshaedcelhkfkhhkfefvgfpgsvkvysynldeqctkfypdpdktgvin	1491
	I : : :: : : : : : : : : : : :	
Qy	1460 VILQLNSIPSSDFLCVRFRIFELFEVGFGLSPATFTYYEHRPDKQCTMEYSTSNIK-I-Q	1517
Db	1492 kichgnicrcaetcsllnqq-k-kidlqlriqkacagndvyvktklrlieekdgndiy	1549
	I : : : : : : : : : : : : : : :	
Qy	1518 KVEGAACKCEADCCQEQEELDLTISAEIRKQTACKQEIAAYAKVISITSITWENFVKY	1577
Db	1550 fmdvlviktgdtnaqakarqvsgkrqcealnklhdnylwlglesdlwpmk-d-dis	1607
	: : : : : :: : : : : : : : : : : :	
Qy	1578 KATLLDDTYKTGEAVAEKDEITFIKKVTCTNA-ELVGQRQYLING-KEAL-OIKYNFSFR	1634
Db	1608 ylit-kn-twierwpnedecdq-eef-qmlcddfaq	1639
	I : : : : : : : : : : : :	
Qy	1635 YIYPLDSLTIWEYWRPDTCSSQCAFANL-DEFAE	1669
 RESULT 3		
ID	R94030 standard; Protein; 1663 AA.	
AC	R94030;	
DT	21-MAY-1996 (first entry)	
KW	Human modified C3 (D752G, E753S, D754G).	
DE	C3 protein; complement; Factor I; Factor H; complement; tumour;	
KW	infection; therapy.	
OS	Synthetic.	
FH	Key Location/Qualifiers	
FT	Peptide 1..22	
FT	/label= Sig_peptide	
FT	Protein 23..667	
FT	/note= "C3 beta chain"	
FT	Peptide 668..671	
FT	/note= "amino acids 668-671 are removed when the	
FT	precursor is cleaved into the alpha and	
FT	beta chains"	
FT	Protein 672..1663	
FT	/note= "C3 alpha chain"	
PN	WO9607738-A2..	
PD	P14-MAR-1996.	
PF	08-SEP-1995; G02121.	
PR	08-SEP-1994; GB-018147.	
PR	04-MAY-1995; GB-009102.	
PI	(IMUT-) IMUTRAN LTD.	
PA	Farries TC, Harrison RA;	
DR	WPI; 96-171613/17.	
PT	Mutant complement pathway protein forming stable C3 convertase -	
PT	for generalised complement depletion or localised complement	
PT	activation	
PS	Claim 11; Fig 1; 81pp; English.	
CC	A modified human C3 protein (R94030) differs from the wild-type	
CC	(R94028) by substitution of Asp-Glu-Asp at positions 752-754 by	
CC	Gly-Ser-Gly. It is obtained by site-directed mutagenesis of	
CC	C3-encoding cDNA (T17738). The modification reduces the	
CC	interaction of C3b/C3i with Factor H in comparison to wild-type	
CC	C3. This allows the modified C3 to be used therapeutically to	
CC	super-active the complement system or the increase a target's	
CC	(e.g. tumour, pathogen or virus-infected cell) sensitivity to	
CC	complement-mediated destruction.	
SQ	Sequence 1663 AA;	
 Query Match 19.4%; Score 2339; DB 16; Length 1663;		
Best Local Similarity 28.3%; Pred. No. 1.02e-194;		

FT /label= Sig_peptide
FT Protein 23..1667
FT /note= "C3 beta chain"
FT Peptide 668..671
FT /note= "amino acids 668-671 are removed when the
FT precursor is cleaved into the alpha and
FT beta chains"
FT Protein 672..1663
FT /note= "C3 alpha chain"
FN #09607738-A2.
PD 14-MAR-1996.
PR 08-SEP-1995; G02121.
PR 08-SEP-1994; GB-018147.
PR 04-MAY-1995; GB-009102.
PA (IMUT-) IMUTRAN LTD.
PI Farries TC, Harrison RA;
DR WPI; 96-171613/17.
DR N-PSDB; T17738.
DR Mutant complemented pathway protein forming stable C3 convertase -
PT for generalised complement depletion or localised complement
PT activation
PS Disclosure; Fig 1; 81pp; English.
CC Human C3 protein (R94028) was produced by expression of a cDNA
CC sequence (T17738) isolated from a human liver cDNA library.
CC C3 is a complement pathway protein that is acceptable to cleavage
CC by Factor I and is also susceptible to the inhibitory action
CC of Factor H. Mutants of C3 (R94029 and R94030) have been
CC produced by site-directed mutagenesis. These mutants can be
CC used to super-activate the complement system, or to induce
CC localised super-activation at a specific target to increase
CC the target's sensitivity to complement-mediated destruction.
SQ Sequence 1663 AA;

Query Match 19.4%; Score 2339; DB 16; Length 1663;
Best local Similarity 28.3%; Pred. No. 1.02e-194;
Matches 488; Conservative 458; Mismatches 661; Indels 115; Gaps 90;

Db 8 sll-ll-llthlpalag-spmyisilpnllrseetmyleahdagqdvptvtvhdffg 64
Qy 2 GLLGILCFLEFLKQWGEQYTWISAPKIFRVGASENIVIQVGYTEAFDAVISIKSYD 61

Db 65 kklvlseektvlpnthmgnvftipaneefkeekgrnkfvvtqatftqgve-kwlv 123
Qy 62 KKFYSYSGHVHLSSENKFNQNSAIIITQIP-KQLPGGQNPVSYVYLEVV-SKHFPSKRWPI 119

Db 124 slsgyflqtdktytpgstvlrftvnhkllpvgtrvmvniempgipvkqdslesq 183
Qy 120 TYDNGELEIHTDKPVYTPDQSKVRVYSINDDLKPAKRETVLFTIDPEGEV--DWVEI 177

Db 184 nqlvlples-wdipelmvgqkwirayenspqqvfstefevkeylvlpsefivteptkf 242
Qy 178 DHGIISPPDFKIPSNRYGMMWIKAKYKEDFSTGTAYEEVKEVLPVHFVSIEP-E-Y 235

Db 243 yyl-yne-kglevtitarflygkkv-egtatvifgi-qd--geqrialspeelkripiedq 296
Qy 236 NFICYKFNQNFETIKARYFYFNWVTEADVITFGIREDIJDKQKEMQMTAMQNTMLNG 295

Db 297 sgevlrskvllldgvpnaedilvgkelysatevilhsgsdmvaqersipivtspqih 356
Qy 296 IAQVTFDSETAVKELSYSLIEDLNKKYLYIAVTVIESGTGSEAEIPGKYKVLSPYKLN 355

Db 337 ftktkyfkpompfdlmvfnvp-d--gs-pay-rvp-vav-q--ge-dtvgaltq-qd 404
Qy 356 LVATPLFLKPGIPPIKQVKQSDLDLUGGVPIVLAQITIDVNGETSDLPKSVTRVDD 415

Db 405 qvaklsinthpsqkplsvtrtkkqelseaeatrtmqalpystvqnsnnylnhlevlrte 464
Qy 416 GVASFVINLPISGVTVLEFNVTADPDLPEENOAREGYRAIYSSLSQSYLYIDWTDNIKA 475

Db 465 lrpetlnvfillrmdraheakirytyliimnkrlkagrqvrepqgdllvplaittd 524
Qy 476 LLVGEHLAI-IVTPKS-PYIDKITHYILILSKGKIHFCTREXESDASYSQISINPVTON 533

Db 525 fipefrlvayvtlligasqrevadsvvvdvdkdcvqslvsksgsedrqpvpqgmtilk 584
Qy 534 MPTSRLIVYIVTGEQTA-ELVSDSVMLNTEEXCGNLQVHLSPADAYS-PCQTVSLN 591

Db 585 iegdhgarvliavdgfvlnknkltqekiwdvvekadictpggkdyagvsdagl 644
Qy 592 MATGMDSVVALAAVDSAVYGVQRCAGKPLEARVQFLEKSDLCGGGAGGLNNANVPHLAGL 651

Db 645 tftssgqgtatraelqcpqaarrtrsvqltekrmdkvq-kypkel-rkccedgmtenp 702
Qy 652 TFLTNANADDSQENDEPCKE-ILPRRT--L-QKKIEETAAKYKHSVVKCCYDGCACVNN 707

Db 703 mrfscqrtrfislgeackvfldeccnyitelrrqharashlglaranlidediaeeniv 762
Qy 708 DE-TCQRAARISIGPRCTKAFTECCVWASQIRANISH-KDMQLGRUHK-TLLPVSKE 764

Db 763 srsefpeawlnvedikeppkngistklmnlfldsittweilavmsdkkgicvadpfe 822
Qy 765 IRSYFPESWLMEVH-L-VPRRK----QL-QFALPDSLTTWEIOGIGISNT-GICVADTVK 816

Db 823 vtvmqdfidrlpsvsvrneqveiravlynrgqlkvrvellhnpafcslatktrh 882
Qy 817 AKVKDFVLEWNTIPYSVVRGEQIQKCTVYNYRTS-QMQFCVKMSAVEGICITESPYIDH 875

Db 883 qqtit---ippk--sals--vpyvivplktglqevekaavyhhfisdgvrkslkvypge 935
Qy 876 QGTCKSKVORQKVEGSSSHLVTFVLPLEIGHNINFSLETW--FGKEILVKTILVWVPE 933

Db 936 irmnktavrtldperl-gregvqkedip-padlsdqvdptesetril-lqgtpvagmte 992
Qy 934 VKRESYGV-TLDPFGIYGTISRKE-FPYRIPL-DLVPKTEI-KRILSVKGLLVGEILS 989

Db 993 daydaerlkhliutpgcgeqgmigmtptviavhyldeteqwekf-q--lekrqgaleli 1049
Qy 990 AVLSQEGINI LTHLPKGSAAELMSVVPVYVYHYLETGNHWNIFHSDPLIEKQKIKKKL 1049

Db 1050 kkgytqqlafrcpsaafaafkraptwtlayvyvkslavnlaidsvlcgvakwlll 1109
Qy 1050 KGLMSLSYENADYSYVWKGSASTWLTAFALRVILGQVNYVEQNQNSICNSLJLWVE 1109

Db 1110 ekqtkpdyfgedapvihgemiqlr-nnnekdmaltafvlsalgeakdiceeqvnslpgs 1168
Qy 1110 NYQUNGSKFENSOYQPIKIAQGTLPVEARENSLYLTAFTVIGIRKAFDIP-LVK-IDTA 1167

Db 1169 itkagdflaanymlqrsyvtvaiaqvala--qmgr--lkg--pllnk-flttakdk-nr- 1219
Qy 1168 LTKADNFFLLENTLPAQSTFTLTAISVALSIGDKTHPQFRSIVSAIKREALVKGNPPYRF 1227

Db 1220 w-ed-pgkq--l-y-----veatsyallallqlkdfvppvrvwlneqrvyggvggst 1269
Qy 1228 WKONLQHKDSSVPNTGTARWVETTAVALLTSLNLKIDINYVNPVIKWLSEEQYGGGFYST 1287

Db 1270 qatfmvqalagqkdapdhqelnldvslqlpsrskithrihwesasllreetskeneq 1329
Qy 1288 QDTINAEGLTSL-LVKQLRLSMDIDVSYKHKGALHNYKWT-DKNFLGRPVEVLNDD 1345

Qy 240 YGNFK-NFEITIKARYFNKVVTEADVITFGIREDLKDKQKMMQATQNTMLINGIAQ 298
Db 292 atldrb-frs-rfplnlnelvghtlaavvtmtesgdmvvtteqsgihivaspyqlhftk 349
Qy 299 VTFDSEITAKELSYSLIEDINNYLYIAVTVIESTGSEAEIPGKGYLSPYKLNIVA 358
Db 350 tpykfpmpyeltvvtvnp-d---gs-paa-hvp-v-v-ea--fh-smgtt-ledqta 396
Qy 359 TPLFLKPGIPIPKVQKDSLDLQVGGVPVTLNAQTIDVNOETSDLDPSKSVTRVDGVA 418
Db 397 klilnlnplnagslptvtrnhgdprerqatksmtaiavtqgsgnylhvaiteiekp 456
Qy 419 SFVILPESGVTVLEFNKVTDAOLPEENQAREGYRAIAYSLSQSYLYTDWTDNHRKALV 478
Db 457 gdnlpvfnvknanslkqikyftylilnknkikfkgvqrrdqgnlvmtmhlhitpdilip 516
Qy 479 GERINI-I-VTPKSPYIDKITHNYLILSKGKIHFETREKFSDAYSQISINIPVTQNVVP 536
Db 517 srfvayqv-gnn--eivadsvwvdkvtcmgtl-v-vkgdnliimgpaamkiklegdp 571
Qy 537 SRRLLVYVYTGEOIAELVSDSVNLIEKCGNQLQVHLSPDADAYSPQTVISLNWATCH 596
Db 572 gatvglavdkavylndykieqakidwtiekedfgctagsgmnlgvfedaalatts 631
Qy 597 DSWALANVDSAVYGVQBAKKELEVRQFLEKSDJGCGAGGLNANVFLHAGLFLTN 656
Db 632 tnlnkrgaakcpnarrirrsvllldenaakaefqddlrkccedvmbnpgnyc 691
Qy 657 ANADDQENDEPCKE-ILRPRT---LQKKTEEIAAKYKHSVWKKCCYDAGCVNDE-TC 711
Db 692 ekrazyiqedackaafleccrykygvrdenqreselflarrdnedgfiadsdiarsdf 751
Qy 712 EQRAARISLPRICAKAFTECCVAVSLRA-NISHKDMQLGRL-HMKTLLPVSKPEIRSYF 769
Db 752 pkewlwtldlteepnsggistsmfsfysldsttwr-lavsfptkigicvaepyeirm 811
Qy 770 PESWLEW-HLV--PR-R---KQQLFALPDSLTWEIQIGISNT-GICVADTVKAKVF 820
Db 812 kvffidlmpysvwnegveirailhnyvmedl-y-vr---velly---npafcsastk 863
Qy 821 KDVLENNIPYSVWRGEQIQKGTVTYNTYRSGMQFCVMSAVEGICTSESPIVDHOGTKS 880
Db 864 qryrqfpikalsstavgfvlvplegghdveikaavqealw-edgv-rkklkvpeqvq 921
Qy 881 SKCVRQ-KVEGSSHLVFTPLPLEIGHNINF--SL-ET-WFGREILVKTIRVWPEGVK 935
Db 922 ksvitvklprak-gvgvgtqlvrikarkldrvpdteietkiiigdpdvaqiensidg 980
Qy 936 RESYSGVTLDPRCYICTSRKKE-PPYRIPLDLVLPKTEIKRILSVKGLVGEILSAVLQS 994
Db 981 sklnhlitpscgqeqmirmasapviaty-yldtqwetlginrte-a-vnqvt-gy 1036
Qy 995 EGINILTHLPKSAEAEMLSW-PVFYFHYLETGNHWNIFHSDPLIEKQKLKKLKEGM 1053
Db 1037 aqmqvkkadhsyaaftrassavpivkvfamaakmvagieheicggvrvlllnrq 1096
Qy 1054 LSIHSTRNADYSYVWKGGSASTWLTAFALRVLGQNKYVEQ-NQNSICNSLMLVENVYQ 1112
Db 1097 qpdgafteapvlsgtmagg--iqgaeeeyltafilvalleakticndymldssikk 1154
Qy 1113 LONGSFENSOYQ IKLAGTLPVFARENSLYLTAFTVIGIRKAFDTCF-LVK-IDTALIK 1170
Db 1155 atnyllkkyeklqrpvyttaltayalaadqlndd-r-v--lm-aa-stgrd---h-w-e 1202

Qy 1171 ADFNLTLTLPAGSTFTLAIASVALSIGDKTHPQFSIVSALKREALVKGNPPIYREWKD 1230
Db 1203 --ey-na---ht---hniegtsyallalmmkfdtqgpiwrltdqmfyetygqtat 1253
Qy 1231 NIAHKDSSVPNTGTARWETPAYALLTSLNKDINYNPVINKWLEEQRYGGGFYSTODT 1290
Db 1254 vmafgalaeeygmphthkdlndltielpdevprryinyenallartvetklnqditv 1313
Qy 1291 INAIEGLEFTEYSL-LVKQLRLSMDIDSVYKKGALHNYKWT-DKQFLGRPVEVLNDDLTIV 1348
Db 1314 tas-gdakatniltfynaqlqekancvknkfhlnvsvenihlna-mg-akgalmkiki-ct 1369
Qy 1349 STGFGSLGTLVHTVTVVH-KTSTSEEVCS-FYLIKDTQDIEASHYRGYNSDKRIVACA 1406
Db 1370 ry-lg--evds-tm-tiidiemltgflpdaedlrrlekvgdryisryevdhnmaqvavi 1424
Qy 1407 SYKPSREESSGSHAVMDISLPTGISANEEDIKALIVEGVQDLFTDYQI-KD-GH---VI 1461
Db 1425 iylnkvshsdeclhfkilkhfvgfipgsvkvysynldekctkfyhpdktgglInki 1484
Qy 1462 LQJNSIPSSDFLCVREIRIFELFEVGLSPATFTTYEYHRPDKQCTMEYTSN- IKI-QKV 1519
Db 1485 cignvrcagatcsslnhq-e-rldvplqiekacetnvdvyvktklrlieeqdndiyvm 1542
Qy 1520 CEGACKCEVADCGMQEELDLTISAEHTKQACKEIAYAKYSITSITVENVFVKYA 1579
Db 1543 dvlevikgtqdnprakthqyisqrkcealnlnkvmdvdyliwgsrdlptk-d-kisyi 1600
Qy 1580 TLLDIYKTEAFAEKDSEITFKRVCTNNA-ELVKGQYLLMG-K-EALQIKYNSFRYI 1636
Db 1601 it-kn-twierwphedece-eef-qklcddfaq 1630
Qy 1637 YPLDSUTWIEYWPRTDTCSSQAFANL-DEFAE 1669

RESULT 7

ID R63224 standard; Protein; 1333 AA.
AC R63224;
DT 05-JUL-1995 (first entry)
DE Cobra partial CVF2.
KW Cobra; C3; third component of complement; human; mouse; rat;
KW X. laevis; pre-pro molecule; beta chain; alpha chain; codon usage;
KW immune response; host defence; tumour.
OS Naja naja.
FH Key Location/Qualifiers
FT Region 1..332
FT /note= "Partial alpha chain"
FT Region 416..715
FT /note= "Gamma chain"
FT Region 947..1333
FT /note= "Beta chain"
PN W09423024-A.
PD 13-OCT-1994.
PF 07-APR-1994; U03441.
PR 07-APR-1993; US-043747.
PA (GEOU) UNIV GEORGETOWN.
PI Bredehorst R, Fritzinger DC, Vogel C;
DR WPI; 94-333186/41.
DR N-PSDB; Q77791.
PT DNA encoding cobra C3, CVF 1 and CVF 2 - which are used in the
PT treatment of cancer
PS Claim 5; Fig 2M-2V; 155pp; English.
CC This sequence represents the C-terminal portion of cobra venom factor

RESULT	ID	Query Match	3.6%	Score 429;	DB 10;	Length 344;
AC	R51948 standard; protein; 344 AA.	Best Local Similarity	28.5%;	Pred. No. 1.78e-24;		
AC	R51948;	Matches	101;	Conservative	98;	Mismatches 127; Indels 28; Gaps 17;
DT	08-NOV-1994 (first entry)					
DE	Phospholipase A2 inhibitor - C3dg (Rat).					
KW	Phospholipase A2; Inhibitor; Rat C3dg; production;					
KW	organic solvent; acid solvent; Allergy; Inflammation.					
OS	Rattus sp.					
PN	J0605293-A.					
PD	08-MAR-1994.					
PF	21-AUG-1992; 222693.					
PR	21-AUG-1992; JP-222693.					
PA	(TEIJ) TEIJIN LTD.					
DP	WPI; 94-121221/15.					
PT	Prod. of 14kDaII type phospholipase A2 inhibitor - by treating					
PT	specified proteins with organic and/or acidic solvent, to obtain					
PT	enzyme useful to treat inflammation and allergy					
PS	Claim 1; Page 5-6; 8pp; Japanese.					
CC	A 14kDaII type phospholipase A2 inhibitor, (C3dg) is produced by					
CC	treating the polypeptide R51948 with an organic and/or acid solvent.					
CC	Rat C3dg can be mass-produced by genetic engineering for use in					
CC	the treatment of inflammation and allergic diseases.					
SC	Sequence 344 AA;					
Db	10 dqpddtsetril-lqgtpvagmaedavderlkhliivpsgcgqmigtptviavhy 68					
Qy	966 DLVPRTEI-KRILSVKGLLVEILSAVLSSQEGINILPHLPKGSAAELMSVPVVFVHY 1024					
Db	69 ldgtqewkf-g--lekrqaelelikgvtqqlakfpssayaaafnnrpsbtwtayvkv 125					
Qy	1025 LETGHNHIFSPDPLEIKQKLKKLKEGMLSTMSYRNADYSYVWKGCSASTWLTAFALR 1084					
Db	126 vfslaanliaidsqvlcgavkwilekqkdpdqvfqcdqpvihqemiggr-ntkeadvsl 184					
Qy	1085 VLGVQNVKVFQONSTCNLSLWLVENYQLDNGSKFENSQYQPIKLOGLTPEARENSLYL 1144					
Db	185 tafviallqeardiccgvnslpqsinkagetleasylnlqpytvalagyalalmkle 244					
Qy	1145 TAFVIGIRKAFDIP-LVK-IDTALIKADNFLENLETPAQSTFTLLAISAYLSGDKTH 1202					
Db	245 epy--ltkftint-a-k-drn-r-weepgqq---l-yn-----veatsyallallilk 285					
Qy	1203 POFRSIVSALKREALVKGPPYIRFYKDMQIAHQDSVPNTGTARMVETTAYALLTSINIK 1262					
Db	286 dfdsppvvr-wlneqryvggystgatfmvfgalacyatdvpdhdkldmndvel 339					
Qy	1263 DINYVNPVKWLSIEQRYGGGFTYSTDTINAIEGFTYLSLLVKQLR-LMSWDIDV 1315					

Search completed: Wed Jan 28 12:13:50 1998
Job time : 66 secs.

Query Match	3.7%	Score 450;	DB 4;	Length 344;
Best local Similarity	28.8%;	Pred. No. 2.96e-26;		
Matches	102;	Conservative	99;	Mismatches 125; Indels 28; Gaps 1
bb	10	dqydpdtdetril-lggtpvaaemaadvgerlkhllivtsgcgcgmimgmtptvaviyh	68	
		: : :	: : : :	
yy	966	DLPVPTKEI-KRILSVKGLIVGEILSAVLQSEGINTLTHLPKSGAEALMSVPVFVEHY	1024	
bb	69	ldgtqewkf-g--lekrqealelikkgyttqqafkpgssavaafnnrppstwtlayvvk	125	
		: :: : : :	: : : : : : :	:
yy	1025	LGTGNHNTFHSDP LIEOKKLKKKLKGMLSTMSYRNADYSVSWKGGSGASTWLFAPALR	1084	
bb	126	vfslaaniiaidsvcgavkwllilekpkdgvfqcdgvpvlhqemigfft-ntkeadvsl	184	
		: : : : : : : : :	: : : : : : : :	:
yy	1085	VLGQKNKYEQNQNSISLLNLVNYYLDNGSKFENSQTQPKLQGLTLPVEARENSLYL	1144	
bb	185	tavfualaqeardi cegqvnlpsinkageyleasylnlqrpytvaiaagyvalalmkle	244	
		: : : : : : :	: : : :	:

Query Match	3.68;	Score 436;	DB 3;	Length 344;
Best Local Similarity	28.2%;	Prod. No. 4.57e-25;		
Matches	100;	Conservative 99;	Mismatches 127;	Indels 28; Gaps 17;
bb	10	dvpdtdsetril-lqgtpvqgmadeavdgerlkhliivtpgcgcqgmigmtptviavhy 68		
		: : : : : :		
2y	966	DLVPTKEL-KRILSVKGLLVCEILSAVLQSGEINILTHLPKGSRAEALSVVPVVFYHY 1024		
bb	69	ldqteqekf-g--lekrqealelikgtytqqlafkqesayaafnrrpstwltkayvwk 125		
		: : : : : : : : : : : :		
2y	1025	LETGNHNIIFHSDPLIEKQKLLKKLUKEGMLISMSYNADYSYVWKGSGASTWLTAFALR 1084		
bb	126	vfslaanlaidsqvlqavkwllilektpdqgfyqcdgqvihqemiggfr--ntkeadvsl 184		
		: : : : : : : : : : : : : : :		
2y	1085	VILGQWNVYEONSGISNLILWENYQLDNGSKFKNQYQPKLOQTLPVEARENSLYL 1144		
bb	185	t:fvlliaqearldceqgvnslpgsinkageyleasylnlqrpvtvaiaqyalalmk 244		
		: : : : : : : :		
2y	1145	TAFTVIGIRKAFDICP-LVK--IDTALIKADNFLENTLPAGSTFTLAI SAYALSIGDKTH 1202		
bb	245	epy--ltkflint-a--k--drn--r--weepgqg----ln-----veatsyallalllk 285		
		: : : : : : : : :		
2y	1203	POFRSIVSALKREALVUKGNPPIYRFWKONLQHKDSSVPITGTARWVETTYALLTSINLK 1262		
bb	286	gfdsrvpvrwvneqryvggvgstqatfmvfqaalagyt dvpdhkdalmvsl 339		
		:		
2y	1263	DINYVNPVILKLEEQRYGGGFTYQDITINAEIGTEYSVLVKQLR--LSMDIDV 1315		